

# **Informative Inventory Report Slovenia 2019**



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Submission under the UNECE Convention on Long-Range Transboundary Air Pollution and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants



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- ANNEX 1 Activity data
- ANNEX 2 Inclusion/exclusion of the condensable component from PM<sub>10</sub> and PM<sub>2.5</sub> emission factors
- ANNEX 3 Recommendations from 2018 in-depth EU NECD review

# 1 EXECUTIVE SUMMARY

## 1.1 Background information on emission inventories

This report is Slovenian Annual Emissions Informative Inventory Report (IIR) submitted under the UNECE Convention on Long-Range Transboundary Air Pollution and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants. The report contains information on Slovenian inventories for all years from the base years (1980, 1987 or 1990) of the protocols to the year 2017.

The substances for which there are existing reporting obligations in the Convention and the Protocols include: SO<sub>x</sub> (as SO<sub>2</sub>), NO<sub>x</sub> (as NO<sub>2</sub>), NMVOC, CO, NH<sub>3</sub>, TSP, PM<sub>10</sub> and PM<sub>2.5</sub>, BC, Pb, Cd, Hg, As, Cr, Cu, Ni, Se, Zn, Dioxins/Furans (PCDD/DF), PAHs, HCB, PCB.

Substances for which emission reporting is obligatory:

- SO<sub>x</sub>, which means all sulphur compounds expressed as sulphur dioxide (SO<sub>2</sub>), including sulphur trioxide (SO<sub>3</sub>), sulphuric acid (H<sub>2</sub>SO<sub>4</sub>), and reduced sulphur compounds, such as hydrogen sulphide (H<sub>2</sub>S), mercaptans and dimethyl sulphides, etc.;
- NO<sub>x</sub>, nitrogen oxides, which means nitric oxide and nitrogen dioxide, expressed as nitrogen dioxide (NO<sub>2</sub>);
- NH<sub>3</sub>, ammonia;
- NMVOCs, non-methane volatile organic compounds, which means all organic compounds of an anthropogenic nature, other than methane, that are capable of producing photochemical oxidants by reaction with nitrogen oxides in the presence of sunlight;
- CO, carbon monoxide;
- Particulate matter (PM), which is an air pollutant consisting of a mixture of particles suspended in the air. These particles differ in their physical properties (such as size and shape) and chemical composition. Particulate matter refers to:
  - PM<sub>2.5</sub>, or particles with an aerodynamic diameter equal to or less than 2.5 micrometres (µm)
  - PM<sub>10</sub>, or particles with an aerodynamic diameter equal to or less than 10 µm
- Cadmium (Cd) and its compounds;
- Lead (Pb) and its compounds;
- Mercury (Hg) and its compounds;
- Polycyclic aromatic hydrocarbons (PAHs). For the purposes of emission inventories, the following four indicator compounds shall be used: benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, and indeno(1,2,3-cd)pyrene;
- Dioxins and furans (PCDD/F), which are polychlorinated dibenzo-p-dioxins (PCDD) and polychlorinated dibenzofurans (PCDF), tricyclic, aromatic compounds formed by two benzene rings, connected by two oxygen atoms in PCDD and by one oxygen atom in PCDF, and the hydrogen atoms of which may be replaced by up to eight chlorine atoms;
- Polychlorinated biphenyls (PCBs), which means aromatic compounds formed in such a manner that the hydrogen atoms on the biphenyl molecule (two benzene rings bonded together by a single carbon-carbon bond) may be replaced by up to 10 chlorine atoms;
- Hexachlorobenzene (HCB), Chemical Abstracts Service (CAS) Registry Number 118-74-1.

Substances for which emission reporting is encouraged include:

- Black carbon (BC), which means carbonaceous particulate matter that absorbs light;
- Total suspended particulate matter (TSP);
- Arsenic (As), Chromium (Cr), Copper (Cu), Nickel (Ni), Selenium (Se) and Zinc (Zn) and their compounds.

The annual emission inventory for Slovenia is reported in the new Nomenclature for Reporting (NFR) format as requested in the revised guidelines for reporting emissions and projections data under the Convention LRTAP (ECE/EB.AIR/122/Add.1, decisions 2013/3 and 2013/4). Revised 2014 Reporting guidelines ECE/EB.AIR.125 are adopted for application in 2015 and subsequent years. The guidelines for the implementation of the inventory of air pollutants contain prescribed methods for calculation of emissions, providing a unified framework for reporting and documenting sources for all inventories. One of the main aims of this method is to ensure comparability of data gathered in individual states and that calls for a definition of at least a minimum scope of equal methods, criteria, and estimating procedures.

This report and NFR tables are available to the public on the EIONET central data repository:  
<http://cdr.eionet.europa.eu/si/un/clrtap/>  
[http://cdr.eionet.europa.eu/si/eu/nec\\_revised/](http://cdr.eionet.europa.eu/si/eu/nec_revised/)

## 1.2 National obligations

Slovenia's annual obligations under the UNECE Convention on Long-range Transboundary Air Pollution (CLRTAP) and its Protocols comprising the annual reporting of national emission data on SO<sub>x</sub> (as SO<sub>2</sub>), NO<sub>x</sub> (as NO<sub>2</sub>), NMVOC, NH<sub>3</sub>, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC as well as on the heavy metals (Pb, Cd, Hg, As, Cr, Cu, Ni, Se, Zn) and persistent organic pollutants (PAHs, PCB, Dioxins/Furans and HCB).

Slovenia had succeeded the LRTAP Convention from Yugoslavia in 1992 with the Act on succession notification (OJ of RS - International Contracts No 35/92, 17 July 1992). Protocols that Slovenia ratified under LRTAP Convention are listed below:

- The 1984 Protocol on Long-term Financing of the Cooperative Programme for Monitoring and Evaluation of the Long-range Transmission of Air Pollutants in Europe (EMEP); 41 Parties. Entered into force 28 January 1988 (Slovenia ratified the protocol in 6.7.1992).
- The 1985 Protocol on the Reduction of Sulphur Emissions or their Transboundary Fluxes by at least 30 per cent; 22 Parties. Entered into force 2 September 1987.
- The 1988 Protocol concerning the Control of Nitrogen Oxides or their Transboundary Fluxes; 30 Parties. Entered into force 14 February 1991 (Slovenia ratified the protocol in 5.1.2006).
- The 1991 Protocol concerning the Control of Emissions of Volatile Organic Compounds or their Transboundary Fluxes; 21 Parties. Entered into force 29 September 1997.
- The 1994 Protocol on Further Reduction of Sulphur Emissions; 26 Parties. Entered into force 5 August 1998 (Slovenia ratified the protocol in 7.5.1998).
- The 1998 Protocol on Heavy Metals; 27 Parties. Entered into force on 29 December 2003 (Slovenia ratified the protocol in 9.2.2004).
- The 1998 Protocol on Persistent Organic Pollutants (POPs); 25 Parties. Entered into force on 23 October 2003 (Slovenia ratified the protocol in 15.11.2005).



- The 1999 Protocol to Abate Acidification, Eutrophication and Ground-level Ozone; 20 Parties. Entered into force on 17 May 2005 - Gothenburg Protocol. Guidance documents to Protocol adopted by decision 1999/1 (Slovenia ratified the protocol in 4.5.2004).

Slovenia has also obligations under European legislation, under the DIRECTIVE (EU) 2016/2284 of the European Parliament and of the Council of 14 December 2016 on the reduction of national emissions of certain atmospheric pollutants, amending Directive 2003/35/EC and repealing Directive 2001/81/EC.

The new Directive repeals and replaces Directive 2001/81/EC, the National Emission Ceilings Directive (NEC Directive) from the date of its transposition (30 June 2018) ensuring that the emission ceilings for 2010 set in that Directive shall apply until 2020. Directive 2016/2284 also transposes the reduction commitments for 2020 taken by the EU and its Member States under the revised Gothenburg Protocol and sets more ambitious reduction commitments for 2030 so as to cut the health impacts of air pollution by half compared with 2005.

Slovenia has obligations under the Regulation (EC) No 850/2004 of the European Parliament and of the Council of 29 April 2004 on persistent organic pollutants (POPs) and amending Directive 79/117/EEC.

### **1.3 Responsible organization**

Slovenian Environment Agency (SEA) is responsible for the annual preparation and submission to the UNECE-LRTAP Convention and European Commission of the annual Slovenian emissions report and the inventories in the NFR format in accordance with the guidelines. Slovenian Environment Agency is independent part of Ministry of the Environment and Spatial Planning.

Slovenian Environment Agency participates in meetings under the UNECE Task Force on Emission Inventories and Projections and the related expert panels, where parties to the convention prepare the guidelines and methodologies on inventories.

### **1.4 Emission trends**

#### **1.4.1 Emission trends for main pollutants**

The main part of the SO<sub>x</sub> emission originates from combustion of fossil fuels, mainly coal and oil in public power plants and district heating plants. From 1980 to 2017, the total emission decreased by 98 %. The large reduction is largely due to installation of desulphurisation plant, use of fuels with lower content of sulphur in public power and district heating plants, introduction of liquid fuels with lower content of sulphur and substitution of high-sulphur solid and liquid fuels to low-sulphur fuels such as natural gas. Despite the large reduction of the SO<sub>x</sub> emissions, these plants make up to 35 % of the total emission. Also emissions from industrial plants, combustion and process emissions are important source of national SO<sub>x</sub>.

The largest sources of emissions of NO<sub>x</sub> are transport followed by combustion in energy industries. The road transport sector is the sector contributing the most to the emission of NO<sub>x</sub> in 2017, 49 % of the Slovenian emissions of NO<sub>x</sub>. The total emissions have decreased by 52 % from 1987 to 2017. The largest reduction of emissions has occurred in power plants and district heating plants due to installation of low-NO<sub>x</sub> burners and denitrifying units. The reductions in

road transport sector have been achieved as a result of fitting three-way catalysts to petrol fuelled vehicles.

Almost all atmospheric emissions of NH<sub>3</sub> result from agricultural activities (91 % in the year 2017). Only a minor part originates from small combustion and transport sector. Road transport sector has been increasing due to increasing use of catalyst cars. The total ammonia emission decreased by 22 % from 1986 to 2017. This is due to decreasing livestock population.

The emissions of NMVOC can be divided into two main groups: incomplete combustion and evaporation. They originate from many different sources. The main contributor of NMVOC in the year 2017 is industrial processes and product use, followed by small combustion. Emissions of NMVOC have decreased from 1990 to 2017 by 54 %. The decline in emissions since 1990 has primarily been due to reductions achieved in the road transport sector due to the introduction of vehicle catalytic converters and carbon canisters on gasoline cars for evaporative emission control, driven by tighter vehicle emission standards, combined with limits on the maximum volatility of petrol as specified in fuel quality directives. The reductions in NMVOC emissions have been enhanced by the switching from petrol to diesel cars and changes in the solvents and product use sector as a result of the introduction of legislative measures limiting the use and emissions of solvents.

CO emissions have decreased between 1980 and 2017 by 67 %. CO is mainly emitted from incomplete combustion. Small combustion is responsible for the dominant share of the total CO emission. Also transport contributes significantly to the total emission of this pollutant. Emission reduction of CO is mainly a result of introduction of vehicle meeting higher emission standards.

#### **1.4.2 Emission trends for persistent organic pollutants (POPs), heavy metals (HM) and particulate matter (PM)**

The persistent organic pollutants and heavy metals emission inventory has been reported for the years 1990-2017.

Persistent Organic Pollutants comprise:

- Polycyclic aromatic hydrocarbons (PAHs):
  - benzo(a)pyrene,
  - benzo(b)fluoranthene,
  - benzo(k)fluoranthene,
  - indeno(1,2,3-cd)pyrene
- Dioxins and furans (PCDD/PCDF or indicated as DF)
- Hexachlorobenzene (HCB)
- Polychlorinated Biphenyls (PCB)

The present emission inventory for PAH (polycyclic aromatic hydrocarbons) includes the four PAHs: benzo(a)pyrene, benzo(b)-fluoranthene, benzo(k)fluoranthene and indeno-(1,2,3-cd)pyrene. The most important source of the PAH emissions is combustion of wood in the residential sector. Small combustion sector contributed 79 % of the total emission in 2017. The PAH emission has decreased by 35 % from 1990 to 2017.

The major part of the dioxins and furans emissions owe to wood combustion in the residential sector, mainly in wood stoves and ovens without flue gas cleaning. Wood and other fuel

combustion in residential plants accounts for 62 % of the national dioxin emission in 2017. Emissions of dioxins and furans have decreased between 1990 and 2017 by 15 %.

The most important source of HCB emissions is electricity and heat production. Among 1990 to 2017 the emission of HCB were decreased by 97 %. The reason for decrease of HCB emissions is termination of HCE use in aluminium production.

Far the most important sources of PCB in Slovenia in 2017 are industrial processes and product use with more than 99 % of the total national emissions. Emissions of PCB were reduced by 91 % in the period 1990 - 2017.

In general, the most important sources of heavy metal emissions are production processes, combustion of fossil fuels and non-industrial combustion and road transport. The heavy metal emissions have decreased substantially in recent years. The reductions span from 98 %, 15 % and 50 % for Pb, Cd and Hg, respectively from the year 1990 to 2017. The reason for the reduced emissions is mainly increased use of gas cleaning devices at power and district heating plants. The large reduction in the Pb emission is due to a gradual shift towards unleaded gasoline, the latter being essential for catalyst cars. Emissions of As, Cr, Cu, Ni, Se, Zn have been estimated for the first time in 2019 submission. Emissions of As, Cr, Ni, Se have decreased by 27, 11, 48, and 32 %, respectively, from the year 1990 to 2017. Emissions of Cu and Zn have increased by 68 % and 17 % between 1990 and 2017.

The particulate matter emission inventory has been reported for the years 2000-2017. The inventory includes the total emission of particles TSP (Total Suspended Particles), emission of particles smaller than 10 µm (PM<sub>10</sub>), emission of particles smaller than 2.5 µm (PM<sub>2.5</sub>) and emissions of black carbon (BC). PM emissions from transport comprise exhaust emissions and non-exhaust emissions from brake and tyre wear and road abrasion. The largest PM<sub>2.5</sub> emission sources are residential plants (73 %). PM<sub>2.5</sub> emissions increased by 6 % from 2000 to 2017. The largest of PM<sub>10</sub> emission sources are also residential plants (66 %) and road transport (10 %). PM<sub>10</sub> emissions have increased by 2 % from 2000 to 2017. The largest TSP emission sources are the residential sector with 59 %. TSP emissions also decreased by almost 1 % from 2000 to 2017. The largest BC emission sources are residential and commercial sector (61 %) and road transport (20 %) as well. BC emissions increased by 2 % from 2000 to 2017. The reason for the increased particulate emissions is mainly due to an increasing wood consumption in residential sector.

## 1.5 General Assessment of Completeness

### Pollutants

SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, NH<sub>3</sub>, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, As, Cr, Cu, Ni, Se, Zn, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, indeno(1,2,3-cd)pyrene, HCB and PCB are covered by the Slovenian inventory.

Emissions of SO<sub>x</sub>, NO<sub>x</sub>, CO have been calculated for the period 1980-2017.

Emissions of NH<sub>3</sub> have been calculated for the period 1986-2017.

Emissions of NMVOC, Pb, Cd, Hg, As, Cr, Cu, Ni, Se, Zn, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, indeno(1,2,3-cd)pyrene, HCB and PCB have been calculated for the period 1990-2017.

Emissions of TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC have been calculated for the period 2000-2017.

### **Geographic coverage**

The geographic coverage is complete. No territory in Slovenia has been left uncovered by the inventory.

### **Notation keys**

#### IE (included elsewhere):

There are a few categories marked with IE in 2017 because relevant data are not available on the reporting level but are included in other category. These sources are:

- 1A3dii National navigation (shipping) – emissions included into 1A3b Road transport
- 1A4aai Commercial/institutional: Mobile – emissions included into 1A3b Road transport
- 1A4bii Residential: Household and gardening (mobile) - emissions included into 1A3b Road transport
- 1A4ci Agriculture/Forestry/Fishing: Stationary - emissions included into 1A4bi Residential: Stationary
- 1A4ciii Agriculture/Forestry/Fishing: National fishing - emissions included into 1A3b Road transport
- 1A5a Other stationary (including military) - emissions included into 1A4ai Commercial/institutional: Stationary
- 2A5c Storage, handling and transport of mineral products - emissions included into 2A1 Cement production, 2A2 Lime production, 2A3 Glass production
- 2C7d Storage, handling and transport of metal products - emissions included into 2C1 Iron and steel production, 2C2 Ferroalloys production, 2C3 Aluminium production, 2C5 Lead production, 2C6 Zinc production, 2C7a Copper production
- 1A3ai(ii) International aviation cruise (civil) - memo items - emissions included into 1A3ai(i) International aviation LTO (civil)
- 1A3aii(ii) Domestic aviation cruise (civil) - memo items - emissions included into 1A3aii(i) Domestic aviation LTO (civil)

#### NE (not estimated):

Notation key NE was applied according to the tables with emission factors in EMEP/EEA Emission Inventory Guidebook, 2016. If in the tables is stated that emission factors for certain pollutants are not estimated, NE was used for particular pollutant and NFR sector.

#### NA (not applicable):

The activity or category exists but relevant emissions and removals are considered never to occur. Application of this notation key is dependent on availability of emission factors in EMEP/EEA Emission Inventory Guidebook, 2016.

#### NO (not occurring)

There are list of sectors marked with NO for the year 2017. NO is used when an activity or process does not exist within a country. No emissions originate from these sectors, since they did not exist in Slovenia in 2017. The highest number of source categories marked with NO is found in agriculture and industrial processes and product use sector, but there are some in

waste and energy industries as well.

-1A1b	Petroleum refining
-1A3di(ii)	International inland waterways
-1A3eii	Other
-1B1b	Fugitive emission from solid fuels: Solid fuel transformation
-1B1c	Other fugitive emissions from solid fuels
-1B2ai	Fugitive emissions oil: Exploration, production, transport
-1B2aiv	Fugitive emissions oil: Refining / storage
-1B2d	Other fugitive emissions from energy production
-2A5a	Quarrying and mining of minerals other than coal
-2A6	Other mineral products (please specify in the IIR)
-2B1	Ammonia production
-2B2	Nitric acid production
-2B3	Adipic acid production
-2B5	Carbide production
-2B7	Soda ash production
-2B10b	Storage, handling and transport of chemical products
-2C2	Ferroalloys production
-2C4	Magnesium production
-2C7b	Nickel production
-2C7c	Other metal production (please specify in the IIR)
-2H3	Other industrial processes
-2J	Production of POPs
-2L	Other production, consumption, storage, transportation or handling of bulk products
-3B4a	Manure management – Buffalo
-3B4f	Manure management - Mules and asses
-3Da2c	Other organic fertilisers applied to soils (including compost)
-3Da4	Crop residues applied to soils
-3Db	Indirect emissions from managed soils
-3Dd	Off-farm storage, handling and transport of bulk agricultural products
-3De	Cultivated crops
-3Df	Use of pesticides
-3F	Field burning of agricultural residues
-3I	Agriculture other
-5B2	Biological treatment of waste - Anaerobic digestion at biogas facilities
-5C1bi	Industrial waste incineration
-5C1biv	Sewage sludge incineration
-5C1bvi	Other waste incineration (please specify in the IIR)
-5C2	Open burning of waste
-5D3	Other wastewater handling
-6A	Other (included in national total for entire territory)
-1A3	Transport (fuel used)
-6B	Other not included in national total of the entire territory (specify in the IIR)
-11A	Volcanoes
-11C	Other natural emissions (please specify in the IIR)

NR (not relevant)

NR is introduced where reporting of emissions is not strictly required by the different protocols. Emission inventory reporting for the main pollutants should cover all years from 1990 onwards if data are available. NR were used for particulate matter before the year 2000.

C (confidential)

Statistical law considering confidentiality is very strict in Slovenia. All data gathered by three or less reporting units is confidential. It is a good practise in national statistic that this boundary is even higher (five units). As Slovenia is a small country, almost all relevant categories from industrial processes sector and, to a lesser extent, from energy sector are also confidential. Nevertheless, no data in our report is marked with C. The confidentiality problem in activity data has been solved on individual level with each relevant plant. After 2005, verified reports from installations included in Emission Trading Scheme (ETS) have resolved this problem generally for most cases.

## 2 INTRODUCTION

### 2.1 Institutional arrangements

In Slovenia, the institution responsible for emission inventories is the Slovenian Environment Agency. In accordance with its tasks and obligations to international institutions, the Slovenian Environment Agency is obligated to perform inventories of GHG and air pollutants emissions within the specified time limit. Slovenian Environment Agency cooperates with numerous other institutions and administrative bodies that relay the necessary activity data and other necessary data for performing inventory each year.

The main source of data is the Statistical Office of the Republic of Slovenia (SORS). Slovenian Environment Agency obtains much of its data through other activities which it performs under the Environmental Protection Act. Emissions from Agriculture are calculated in cooperation with the Slovenian Agriculture Institute. Many data are obtained directly from factories. Inventory institutional arrangements and data sources are presented in Table 2.1.1.

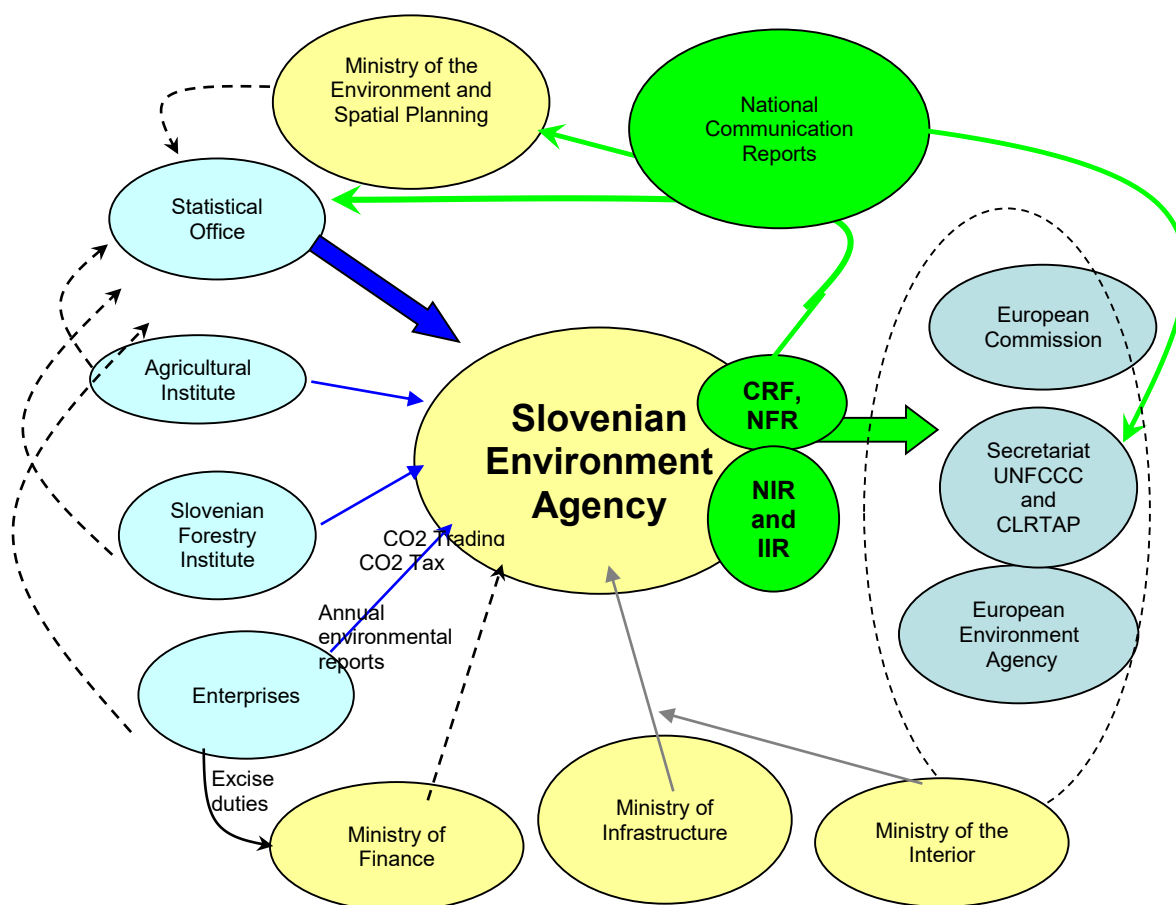
**Table 2.1.1 Inventory Institutional Arrangements and Data Sources**

NFR category	NFR sub-category	Sources of data
<b>NFR 1 A – Energy: Fuel Combustion</b>	NFR 1A1 - Energy Industry	<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia: Joint Questionnaires, Energy Balances, annual energy statistics</li> <li>• Slovenian Environment Agency: ETS data</li> </ul>
	NFR 1A2 - Manufacturing Industries and Construction	<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia: Joint Questionnaires, Energy Balances, annual energy statistics</li> <li>• Slovenian Environment Agency: ETS data</li> </ul>
	NFR 1A3 – Transport	<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Ministry of Infrastructure and Spatial Planning</li> <li>• Slovenian Infrastructure Agency</li> <li>• Slovenian Environment Agency</li> </ul>
	NFR 1A4 – Other Sectors	<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Ministry of the Interior, Police</li> <li>• Ministry of Defence, Slovenian Armed Forces</li> </ul>
<b>NFR 1 B – Energy: Fugitive Emissions from Fuels</b>		<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Slovenian Environment Agency: ETS data</li> </ul>
<b>NFR 2 – Industrial Processes and Product use</b>	NFR 2A – Mineral Products	<ul style="list-style-type: none"> <li>• Slovenian Environment Agency: ETS data</li> <li>• Data obtained from factories</li> </ul>
	NFR 2B – Chemical Industry	<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Slovenian Environment Agency: ETS data</li> <li>• Data obtained from factories</li> </ul>
	NFR 2C – Metal Production	<ul style="list-style-type: none"> <li>• Slovenian Environment Agency: ETS data</li> <li>• Data obtained from factories</li> </ul>
	NFR 2D-2L Other Solvent and Product use	<ul style="list-style-type: none"> <li>• Chemicals Office of the Republic of Slovenia</li> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Slovenian Environment Agency</li> </ul>
<b>NFR 3 – Agriculture</b>		<ul style="list-style-type: none"> <li>• Agricultural Institute of Slovenia</li> <li>• Statistical Office of the Republic of Slovenia</li> </ul>
<b>NFR 5 – Waste</b>		<ul style="list-style-type: none"> <li>• Statistical Office of the Republic of Slovenia</li> <li>• Slovenian Environment Agency</li> <li>• Administration for Civil Protection and Disaster Relief of the Republic of Slovenia</li> </ul>

## 2.2 Brief description of the process of inventory preparation, data collection, processing, data storage and archiving

Owing to the ever-increasing obligations of Slovenia with regard to reporting, the Slovenian Environment Agency has implemented a unified system of data collection for the purposes of making greenhouse gases (GHG) and air pollutants inventories, as well as secures reliable financing in accordance with the annual program of its work.

A Memorandum of Understanding has been concluded with the SORS to submit quality and verified data to the Slovenian Environment Agency in due time, because the time limits for GHG and air pollutants inventories and the national inventory report (NIR) and IIR have shortened with the entry of Slovenia into the EU. In view of this, an agreement has been reached with the participating institutions to shorten the time limits for submitting data. For reasons of complexity, attention was mostly focused on the Joint Questionnaires (JQ) of the SORS, on the basis of which the Statistical Office produces the Energy Balance of the Republic of Slovenia, where in the most important data on the energy sector are to be found. Data flow in the Slovenian Inventory System is presented in Figure 2.2.1.



**Figure 2.2.1 Data flow in the Slovenian Inventory System**

The year 2003 presents the end of the process of harmonization of data collection among the Directorate of Energy, Ministry of Environment and Spatial Planning, and the SORS. An end was put to previous parallel double collecting of data. The competence of collecting data has, by law, passed to the SORS, which checks the data and eliminates potential reporting errors, and submits consolidated data to the Directorate of Energy, which has been publishing data until 2005 in its Energy Yearbook of the Republic of Slovenia. In terms of content, the data were



identical to those submitted in the Joint Questionnaires to the International Energy Agency (IEA).

At the beginning of 2007, the agreement between SORS and the Slovenian Environment Agency came into force. Accordingly, all statistical data which are necessary for preparing emission inventories are available each year by October 30 at the latest. In exchange, European trading scheme (ETS) data and emission estimates are reported to the SORS within a defined time frame. In 2014 the new agreement has been signed which includes more data sets and updated time lines.

A process of inventory preparation is designed according to the PDCA-cycle (Plan – Do – Check – Act). This is a generally accepted model for pursuing a systematic quality work according to international standards, in order to ensure the maintenance and development of the quality system. This structure is in accordance with structures described in decision 19/CMP.1 and in the 2006 IPCC Guidelines. The system consists of inventory planning, inventory preparation, inventory quality checking and follow-up improvements which are integrated into the annual cycle and preparation.

Owing to the ever-increasing obligations of Slovenia with regard to reporting, the Slovenian Environment Agency has decided to implement a unified system of data collection for the purposes of making inventories, as well as secure reliable financing in accordance with the annual program of its work.

For submitting reports to different institutions, various report formats have been devised, since the same data are used to report to the United Nations Framework Convention on Climate Change (UNFCCC), European Environment Agency (EEA), European Commission (EC), and CLRTAP. All external reports of the Slovenian Environment Agency are prepared in accordance with ISO 9001 via the Agency's reporting service, which keeps inventories of reports. Parallel to this, emissions data are submitted to the SORS, which makes this data available in its publications and submits them to EUROSTAT and the IEA.

In 2006, we started to develop a joint database for air pollutants and GHGs. It already contains all activity data, emission factors and other parameters together with a description of sources from 1980 on for other pollutants, and from 1986 on for GHG emissions. At defined control points, QC procedures are included. Some phases of the database were concluded, but the whole process is planned to be finished in 2015. New Nomenclature For Reporting (NFR) and Common Reporting Format (CRF) tables in 2015 required additional changes of the database. Constant improvement of the database is expected.

For each submission, databases and additional tools and submodels are frozen together with the resulting NFR reporting format. This material is placed on central agency's servers, which are subject to routine back-up services. Material which has been backed up is archived safely.

Figure 2.2.1 shows a schematic overview of the process of inventory preparation. The figure illustrates the process of inventory preparation from the first step of collecting external data to the last step, where the reporting schemes are generated for the UNFCCC and EU in the CRF format and to the United Nations Economic Commission for Europe/Cooperative Programme for Monitoring and Evaluation of the Long-range Transmission of Air Pollutants in Europe (CLRTAP - UNECE/EMEP) in the NFR format. For calculations and reporting the software tool is developed by Slovenian Environment Agency.

### 2.3 Brief description of methodologies and data sources used

Slovenia's air emission inventory is based on EMEP/EEA methodology. It has been developed under UNECE/EMEP Task Force on Emission Inventories and Projections (TFEIP) and the European Environment Agency. The basis of inventory is also 2006 IPCC Guidelines for National Greenhouse Gas Inventories. EMEP/EEA (formerly referred as CORINAIR - COoRdination of INformation on AIR emissions) is a European air emission inventory programme for national sector wise emission estimations, harmonized with the IPCC guidelines. To ensure estimates are as timely, consistent, transparent, accurate and comparable as possible, the inventory programme has developed calculation methodologies for most subsectors and software for storage and further data processing. The EMEP/EEA calculation principle is to calculate the emissions as activities multiplied by emission factors. Activities are numbers referring to a specific process generating emissions, while an emission factor is the mass of emissions per unit activity. Information on activities to carry out the EMEP/EEA inventory is largely based on official statistics. The most consistent emission factors have been used, either as national values or default factors proposed by international guidelines. The emission factors used for emission calculations were adopted from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016.

The activity data of consumed fuel energy were provided by SORS. Additional data on the energy use of some types of waste (waste tires, oils and solvents) were acquired from verified ETS reports. Data on fuel consumption in agriculture and forestry refer to mobile sources only, while the rest of the fuel consumption of these sub-sectors is included in the public and service sub-sector. Emissions in road transport were determined with the COPERT 4 model (version 11.4) using default EFs from the model.

Emissions from industrial processes and product use have been mostly determined on the basis of statistical data on production and consumption of raw materials and by applying country-specific emission factors. After 1997, the SORS partly changed the manner of collecting and presenting these data, and therefore most of the data were obtained directly from individual companies (plant communication data) and verified ETS reports.

Important source of data in Industrial processes and product use sector is REMIS database, established and handled by Slovenian Environmental Agency. These data represent plant specific values. REMIS database is obtained in compliance with Rules on initial measurements and operational monitoring of the emission of substances into the atmosphere from the stationary pollution sources and on the conditions for their implementation (OJ RS, No. 105/08). Each year all obligators must provide report on implementation of emission monitoring of substances into air. Annual emission report includes emissions of substances into air. These emissions data are direct measurements of emissions into air and reflect plant specific values.

Additional source of NMVOC data is HOS database. It is similar to REMIS database and it is established and handled by Slovenian Environmental Agency as well. Data in HOS database are obtained in compliance with Decree on limit values for atmospheric emissions of volatile organic compounds from installations using organic solvents (OJ RS, No. 112/05, 37/07, 88/09, 92/10, 51/11, 35/15) and Decree on the emission limit values of halogenated volatile organic compounds into the atmosphere from installations using organic solvents (OJ RS, No. 71/11). Each year all VOC obligators must provide report about solvent management plan (mass balance) for previous year. Data on NMVOC from HOS database have been available since 2005.

Emissions from agriculture and waste sectors have been mostly determined on the basis of statistical data as well. Emission factors used have been mainly obtained from EMEP/EEA Emission Inventory Guidebook, 2016 and by applying country specific emission factors.

Table 2.3.1 Summary report for methods and emission factors used

Categories	Method applied	Emission factors
1. Energy	M,T1,T2,T3	CS,D,M,PS
A. Fuel combustion	M,T1,T2,T3	CS,D,M,PS
1. Energy industries	T1,T2	CS,D,PS
2. Manufacturing industries and construction	T1,T2	D, PS
3. Transport	M,T1,T2	M,CS,D
4. Small combustion and Non-road mobile sources and machinery	T1,T2	CS,D
B. Fugitive emissions from fuels	T1	D,CS
1. Solid fuels	T1	D,CS
2. Oil and natural gas	T1,T2	D
2. Industrial Processes	T1,T2	CS,D
A. Mineral industry	T1,T2	CS,D
B. Chemical industry	T1,T2	CS,D
C. Metal industry	T1,T2	CS,D
D.-L. Other solvent and product use	T1,T2	CS,D
3. Agriculture	T1,T2	CS,D
B. Manure management	T1,T2	CS,D
D. Crop production and agricultural soils	T1,T2	CS,D
5. Waste	T1,T2,D	CS,D
A. Solid waste Disposal on land	T2	D
B. Biological Treatment	T1	D
C. Incineration	T2	D
D. Waste water handling	T1	D
E. Other waste	T1	D

CS - Country Specific, T1 - Tier 1, T2 - Tier 2, T3 - Tier 3, M- Model, D – Default value, PS – plant specific

## 2.4 Key Categories

This chapter presents results of Slovenia's key source analysis. Key categories analysis is increasingly important in order to prioritize emission sources and identify where the implementation of improvements is most effective. We have assessed the most important sources (the sources making up 80% of the national total). The key sources for the 2017 emissions and the corresponding percentages are listed in Table 2.4.1. The analysis of key source categories was performed on the basis of sectorial distribution and using the Tier 1 method and Approach 1. Key categories are those which, when summed together in descending order of magnitude, cumulatively add up to 80 % of the total level.

**Table 2.4.1 List of key sources (and their contribution to total amount) by pollutant for 2017**

Component	Key categories (Sorted from high to low from left to right)											Total (%)
SO <sub>x</sub>	1A1a	2B10a	2C3	1A4bi	1A2f							82,7
	(40,7%)	(13,4%)	(12,6%)	(8,5%)	(7,6%)							
NO <sub>x</sub>	1A3bi	1A1a	1A3biii	1A4cii	1A3bii	1A4bi	1A2f	3Da1				81,6
	(29,2%)	(13,7%)	(13,1%)	(7,1%)	(6,4%)	(5,8%)	(3,2%)	(3,1%)				
NH <sub>3</sub>	3Da2a	3B1b	3B1a	3Da1	1A4bi							80,8
	(41,0%)	(14,2%)	(11,5%)	(7,5%)	(6,6%)							
NMVOC	1A4bi	3B1b	2D3d	2D3a	2D3g	3B1a	1A4cii	1A3bi	1B1a	2H2	1A2gviii	82,2
	(22,7%)	(10,3%)	(8,8%)	(8,3%)	(7,6%)	(6,6%)	(5,0%)	(3,7%)	(3,6%)	(3,0%)	(2,5%)	
CO	1A4bi	1A3bi	2C3									84,7
	(64,2%)	(13,0%)	(7,4%)									
TSP	1A4bi	2A2	1A3bvi	1A3bvii	1A1a	1A3bi	3B4gi	1A2gviii				81,2
	(58,9%)	(4,6%)	(4,2%)	(3,6%)	(3,1%)	(2,5%)	(2,2%)	(2,1%)				
PM <sub>10</sub>	1A4bi	1A3bvi	1A1a	1A3bi	1A2gviii	2A2	1A3bvii					82,1
	(65,8%)	(3,7%)	(3,1%)	(2,9%)	(2,3%)	(2,1%)	(2,1%)					
PM <sub>2,5</sub>	1A4bi	1A3bi	1A1a	1A2gviii								81,4
	(72,8%)	(3,3%)	(2,8%)	(2,5%)								
Pb	1A3bi	2C1	1A1a	1A3bvi	1A4bi							85,9
	(35,8%)	(24,5%)	(9,6%)	(8,9%)	(7,2%)							
Hg	1A1a	2C1	5C1bv	1A4bi	2D3a	1A2d	5C1biii					85,1
	(21,5%)	(20,7%)	(15,5%)	(7,3%)	(7,1%)	(6,7%)	(6,5%)					
Cd	1A4bi	2C1	1A1a	1A2gviii								82,3
	(41,7%)	(22,7%)	(13,5%)	(4,4%)								
DIOX	1A4bi	2C1	5E									84,6
	(62,1%)	(13,1%)	(9,4%)									
PAH	1A4bi	2C1										85,0
	(78,8%)	(6,2%)										
HCB	1A1a	1A4bi	1A2f									87,1
	(61,6%)	(17,9%)	(7,5%)									

## 2.5 Quality assurance, quality control and verification plan

In 2014, Slovenia developed and implemented a Quality Assurance and Quality Control plan. At the end of 2013, a QA/QC manager at the inventory agency was designated. It has been commonly used in preparation of GHG and air pollutant inventories.

Quality Control (QC) is a system of routine technical activities to measure and control the quality of the inventory as it is being developed. The QC system is designed to:

- provide routine and consistent checks to ensure data integrity, correctness and completeness;
- identify and address errors and omissions;
- document and archive inventory material and record all QC activities.

The final part of this system is incorporated in an Oracle database (ISEE – "Emission inventory" information system). ISEE enables and ensures that all necessary built-in QA/QC checks have been performed before data and emission estimates are entered in the reporting format tables. It also keeps a record of all changes made to data in the database.

As all calculations are performed in the database with software generated for this purpose, no human errors are expected. But for QA/QC purpose all emissions are also calculated in the old way in Excel spreadsheets. Both estimates were then compared and all differences were carefully investigated and corrected.

The main purpose of ISEE is:

- to enable collection and archiving of activity data, emission factors and other parameters including descriptions of sources from 1980 on for air pollutants, and from 1986 on for GHG emissions,
- to calculate GHG and air pollutants emissions,
- to automatically fill in reporting tables.

During development of the database, the following QC was performed:

### Check of methodological and data changes resulting in recalculations

- check for temporal consistency in time series input data for each source category
- check for consistency in the algorithm/method used for calculations throughout the time series.

### Completeness checks

- confirm that estimates are reported for all source categories and for all years from the appropriate base year to the period of the current inventory
- check that known data gaps that result in incomplete source category emissions estimates are documented
- compare estimates to previous estimates: for each source category, current inventory estimates should be compared to previous estimates. If there are significant changes or departures from expected trends, recheck estimates and explain any differences.

### Check of activity data, emission factors and other parameters

- cross-check all input data from each source category for transcription errors
- check that units are properly labelled in calculation sheets
- check that units are correctly carried through from beginning to end in calculations
- check that conversion factors are correct
- check that temporal and spatial adjustment factors are used correctly.

#### Check of emissions estimates

For the entire period 1980–2016, emissions are also calculated in the old way using Excel spreadsheets and in the database using built-in formulas. Both estimates were compared and all differences carefully investigated.

The reasons for differences were the following:

- formulas for calculation of emissions were not correct
- data field was not properly labelled
- data relationship was not correct
- emissions data were not correctly aggregated from lower reporting levels to higher reporting levels.

All errors were corrected and the accuracy of emissions calculations on all levels is now assured.

QA/QC checks not performed in the database:

#### Preparation of IIR

- check that all chapters from annotated IIR are included in the IIR
- check that AD, EF and other numerical information mentioned in the text is correct
- check all AD data is presented in the tables in the IIR
- check all EF and other parameters used in the tables in the IIR
- check all graphs for accuracy and presence in the whole period
- check all titles for tables and pictures
- check that all Annexes to the IIR are included and updated

#### Documentation and archiving

All inventory data are now stored in a joint database. Supporting data and references are stored in electronic form and/or hard copy form. Inventory submissions are stored mostly in electronic form at various locations and on various media (network server, random-access memory, computer hard disk). Access to files is limited in accordance with the security policy. Backup copies on the server are made at regular intervals in accordance with the requirements of the information system. All relevant data from external institutions are also stored at the SEA.

QA/QC checks of documentation and archiving procedures:

- check that inventory data, supporting data and inventory records are archived and stored to facilitate detailed review
- check that all supporting documentation on QA/QC procedures is archived
- check that results of QC analysis and uncertainty estimates are archived
- check that there is detailed internal documentation to support the estimates and enable duplication of emissions estimates.
- check that documentation of the database is adequate and archived.
- check that bibliographical data references are properly cited in the internal documentation and archived.
- check that inventory improvements plan is updated and archived.

In 2006, an additional quality control check point was introduced by forwarding the assessment of verified emission reports from installations included in the National Allocation Plan to the SORS. The role of SORS is to compare data from installations included in the EU-ETS with data from their reporting system and to propose corrective measures, if necessary. The outcome of data consistency checks is used as preliminary information for the Ministry of the Environment and Spatial Planning to perform on-site inspections. The use of (EU) ETS data is described in more detail in the relevant chapter on Energy and Industrial Processes sectors.

### Quality assurance (QA)

Quality assurance generally consists of independent third-party review activities to ensure that the inventory represents the best possible estimates of emissions and removals, and to support the effectiveness of the QC program. In the past we have performed only one peer review. In 2006, we received many useful comments from the team preparing our fourth National Communication Report. Although the comments were not presented as an official report, we accepted many of the suggestions and corrected a number of errors. We are planning a sectorial review of our inventory on a yearly basis – one sector per year. In May 2009, a peer review of the Slovenian inventory was performed for the energy sector.

SORS is our main data provider. In 2005, the European Statistics Code of Practice was adopted, bringing considerable changes to the SORS QA/QC system. The main pillars (factors) of quality are defined and thoroughly described in the Medium-term Programme of Statistical Surveys 2008–2012 (<http://www.stat.si/doc/drzstat/SPSR-ang.pdf>). The strategic directions from the Medium-term Programme of Statistical Surveys are presented in detail at [http://www.stat.si/doc/drzstat/kakovost/TQMStrategy\\_2006\\_eng.doc](http://www.stat.si/doc/drzstat/kakovost/TQMStrategy_2006_eng.doc) in the Total Quality Management Strategy 2006–2008.

### **Official consideration and approval of the inventory**

Before the inventory is reported to the EU, EEA, CLRTAP or UNFCCC Secretariat, it goes through an approval process. The institution designated for approval is the Ministry of the Environment and Spatial Planning.

### **Public Availability of the Inventory**

The inventories are publically available on the web. Every submission is accompanied with a short description in Slovenian language. The estimates are presented in a more simple way suitable for general public. Air pollutant emissions are also presented as indicators.

Web page address:

[http://okolje.arso.gov.si/onesnazevanje\\_zraka/vsebine/onesnazevala-zraka](http://okolje.arso.gov.si/onesnazevanje_zraka/vsebine/onesnazevala-zraka)

## 2.6 Description and interpretation of emission trends by gas

### 2.6.1 Emission Trends for Main Pollutants

Emission trends for main pollutants (SO<sub>x</sub>, NO<sub>x</sub>, NH<sub>3</sub>, NMVOC and CO) from years 1980 for SO<sub>x</sub>, NO<sub>x</sub>, CO, 1986 for NH<sub>3</sub> and 1990 for NMVOC to 2017 are represented in Table 2.6.1.1. Emissions decreases are: SO<sub>x</sub> (98 %), NO<sub>x</sub> (49 %), NH<sub>3</sub> (22 %), NMVOC (54 %) and CO (67 %). Target values for the year 2010 and later are for SO<sub>x</sub> (27 kt), NO<sub>x</sub> (45 kt), NH<sub>3</sub> (20 kt) and NMVOC (40 kt).

**Table 2.6.1.1 National total emissions, emission trends and emission target for the year 2017**

Year	Emissions (kt)				
	SO <sub>x</sub>	NO <sub>x</sub>	NH <sub>3</sub>	NMVOC	CO
1980	237,4	67,8			319,8
1981	257,9	68,2			306,1
1982	259,6	66,4			290,1
1983	274,3	64,7			269,4
1984	252,7	64,4			281,4
1985	242,9	65,1			300,3
1986	249,6	70,9	24,0		321,6
1987	229,6	72,0	24,0		331,9
1988	217,5	72,1	23,5		314,9
1989	219,0	71,9	23,0		311,7
1990	201,1	71,0	22,9	64,2	304,6
1991	186,4	65,2	21,6	61,6	285,6
1992	193,6	65,7	22,4	60,4	275,3
1993	190,5	69,5	20,8	61,0	289,2
1994	184,1	72,4	20,8	62,4	282,1
1995	124,0	71,4	20,9	62,3	280,9
1996	115,5	74,1	20,4	65,9	289,6
1997	119,1	72,8	20,7	62,1	261,7
1998	109,9	65,8	20,8	57,3	225,2
1999	96,0	59,2	20,7	54,0	203,0
2000	93,6	58,6	21,6	52,5	187,7
2001	63,1	58,5	21,4	51,6	181,8
2002	62,7	58,0	22,5	50,5	178,1
2003	60,2	55,5	21,4	49,7	176,8
2004	50,7	53,9	19,8	47,7	165,9
2005	40,4	55,2	20,4	44,6	162,6
2006	17,1	55,6	20,4	44,9	153,9
2007	15,6	53,9	21,0	43,1	147,3
2008	14,6	58,1	19,7	41,9	143,1
2009	11,9	49,7	20,3	38,7	131,8
2010	10,7	48,3	19,7	37,3	130,3
2011	13,2	47,5	18,9	34,9	128,0
2012	11,6	46,0	18,7	33,1	122,8
2013	13,8	44,0	18,5	32,5	121,7
2014	9,8	39,1	18,5	29,8	101,7
2015	5,4	35,1	18,7	29,8	107,5
2016	4,6	36,0	19,0	30,1	110,0
2017	4,9	34,7	18,6	29,8	105,0
<b>Reduction trend (%)</b>	<b>-97,9 %</b>	<b>-48,8 %</b>	<b>-22,4 %</b>	<b>-53,6 %</b>	<b>-67,2 %</b>



## SO<sub>x</sub> Emissions

National SO<sub>x</sub> emissions steadily decreased from the year 1980, when total amount was 237,4 kt to 4,9 kt in 2017. Emissions have decreased by 97,9 % between 1980 and 2017. The reduction in emissions since 1980 has been achieved as a result of a combination of measures, including fuel-switching in energy-related sectors away from high-sulphur solid and liquid fuels to low-sulphur fuels such as natural gas, the fitting of flue gas desulphurisation abatement technology in thermal power plants and industrial facilities and the impact of European Union directives relating to the sulphur content of certain liquid fuels.

The highest drop of emission was occurred in electricity and heat production. Important factor of lower emissions from thermal power plants was introduction of flue gas desulphurization device and gas turbines in power cogeneration plants. In 1995, SO<sub>2</sub> emissions fell considerably, mostly due to the operation of the device for the desulphurization of flue gases in unit 4 of the Šoštanj Thermal Power Plant. In the 2001 and 2005, SO<sub>2</sub> emissions again fell considerably, due to the operation of the device for the desulphurization of flue gases (FGD) in unit 5 of the Šoštanj Thermal Power Plant (2001) and Thermal Power Plant Trbovlje (2005).

The 2010 national emission ceiling for SO<sub>x</sub> in Slovenia is 27 kt regarding Gothenburg Protocol and DIRECTIVE 2001/81/EC of the European Parliament and of the Council of 23 October 2001 on national emission ceilings for certain atmospheric pollutants. Slovenia has reduced national SO<sub>x</sub> emissions below the level of the 2010. Total emissions of SO<sub>x</sub> were in the year 2017, 82 % below the national emission ceiling.

The 2012 revision of the Gothenburg Protocol to the UNECE LRTAP Convention and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants set emission reduction targets for SO<sub>x</sub> based on 2005 emission totals, to be met by countries in or before 2020. Reduction of emissions has to be 63 % compared to 2005 emissions. Emissions for Slovenia in 2017 were below a linear target path to its 2020 target by 88 % of its 2005 emission totals.

Slovenia in 2017 fulfilled all requirements under 2nd Sulphur Protocol.

Sulphur dioxide is emitted when fuels containing sulphur are combusted. It is a pollutant which contributes to acid deposition which in turn can lead to changes occurring in soil and water quality. The subsequent impacts of acid deposition can be significant, including adverse effects on aquatic ecosystems in rivers and lakes and damage to forests, crops and other vegetation. SO<sub>x</sub> emissions also aggravate asthma conditions and can reduce lung function and inflame the respiratory tract, and contribute as a secondary particulate pollutant to formation of particulate matter in the atmosphere, an important air pollutant in terms of its adverse impact on human health. Further, the formation of sulphate particles in the atmosphere after their release results in reflection of solar radiation, which leads to net cooling of the atmosphere.

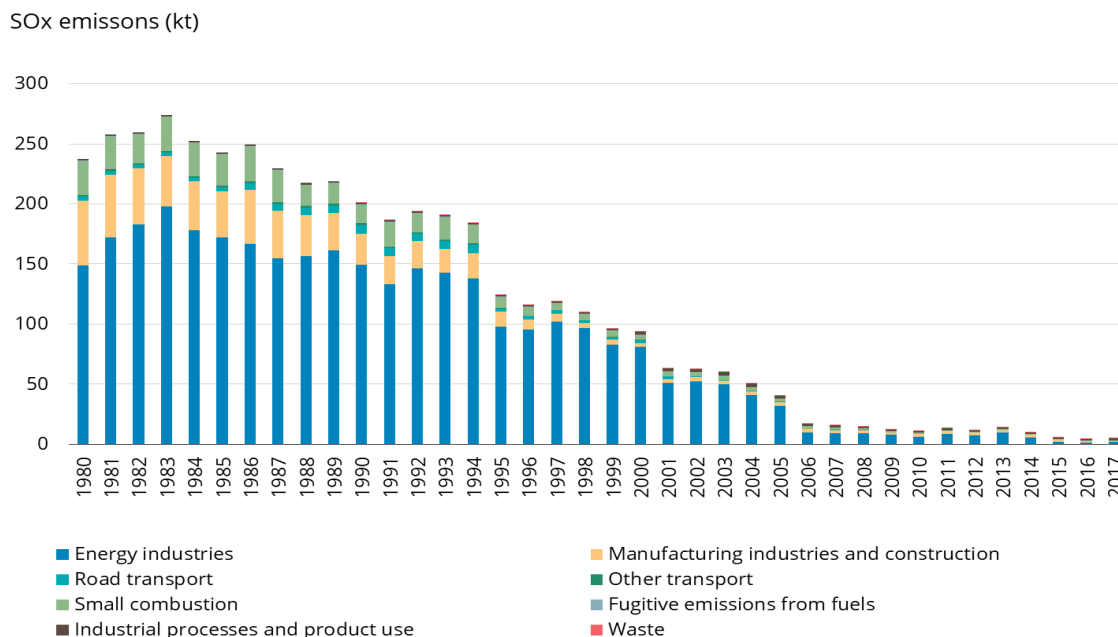


Figure 2.6.1.1 SO<sub>x</sub> emissions in Slovenia for the period 1980 – 2017

In 2017, the most significant sector source of SO<sub>x</sub> emissions was energy industries (41 % of total emissions), followed by emissions occurring in the industrial processes and product use (31 %) and from manufacturing industries and construction (16 %).

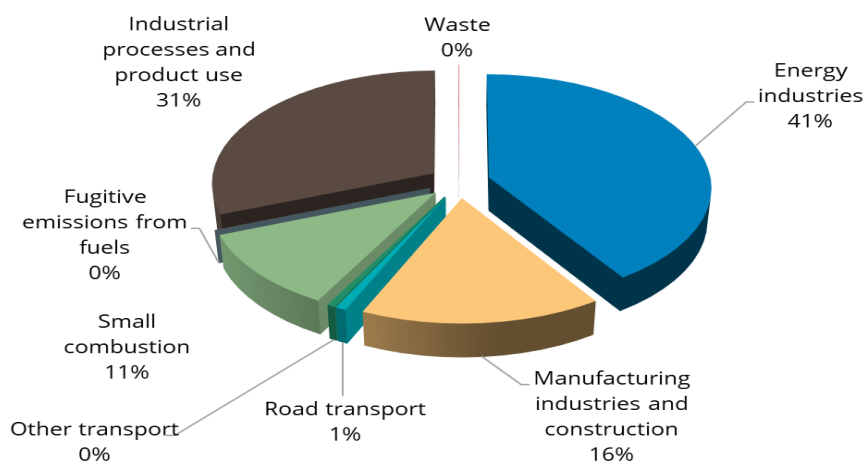


Figure 2.6.1.2 Individual sectors contribution of SO<sub>x</sub> emissions for 2017

## **NO<sub>x</sub> Emissions**

Total national NO<sub>x</sub> emissions in Slovenia decreased from 72,0 kt in 1987 to 34,7 kt in the year 2017. Emissions were reduced by 51,8 %. Despite the base year for NO<sub>x</sub> is 1987 emissions have been calculated from 1980 onwards due to availability of activity data for the whole period. Emissions were reduced by 48,8 % in the period 1980-2017. The largest reduction of emissions since 1980 has occurred in the electricity/energy production sector as a result of measures such as the introduction of combustion modification technologies (such as use of low NO<sub>x</sub> burners), implementation of flue-gas abatement techniques (NO<sub>x</sub> scrubbers and selective catalytic and non-catalytic reduction techniques) and fuel-switching from coal to gas. These reductions have been achieved also in the road transport sector despite the general increase in activity within this sector since the early 1990s and have primarily been achieved as a result of fitting three-way catalyts to petrol fuelled vehicles.

Target value for NO<sub>x</sub> according to Gothenburg Protocol and DIRECTIVE 2001/81/EC of the European Parliament and of the Council of 23 October 2001 on national emission ceilings for certain atmospheric pollutants for year 2010 is 45 kt NO<sub>x</sub>. Slovenia met that target value in 2017, emissions were 23 % below national ceiling value.

The 2012 revision of the Gothenburg Protocol to the UNECE LRTAP Convention and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants set emission reduction targets for NO<sub>x</sub> based on 2005 emission totals, to be met by countries in or before 2020. Reduction of emissions has to be 39 % compared to 2005 emissions. Emissions for Slovenia in 2017 were below a linear target path to its 2020 target by 37 % of its 2005 emission totals. Additional measures may therefore need to be undertaken in future years to achieve reduction target implied by the protocol.

Slovenia in 2017 fulfilled requirements under NO<sub>x</sub> Protocol.

NO<sub>x</sub> contributes to acid deposition and eutrophication of soil and water. The subsequent impacts of acid deposition can be significant, including adverse effects on aquatic ecosystems in rivers and lakes and damage to forests, crops and other vegetation. Eutrophication can lead to severe reductions in water quality with subsequent impacts including decreased biodiversity, changes in species composition and dominance, and toxicity effects. NO<sub>x</sub> is associated with adverse effects on human health, as at high concentrations it can cause inflammation of the airways and reduced lung function, increasing susceptibility to respiratory infection. It also contributes to the formation of secondary particulate aerosols and tropospheric ozone in the atmosphere, both of which are important air pollutants due to their adverse impacts on human health and other climate effects.

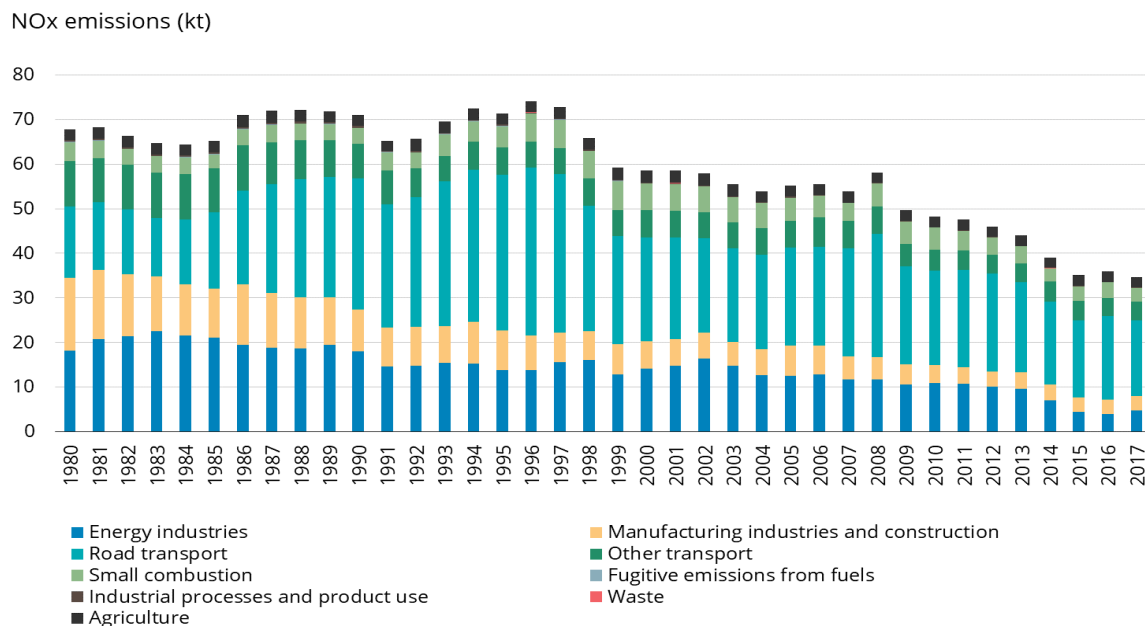


Figure 2.6.1.3 NO<sub>x</sub> emissions in Slovenia for the period 1980 - 2017

In 2017, the most significant sources of NO<sub>x</sub> emissions were the road transport (49 %), other transport sectors (12 %) and energy production and distribution (14 %).

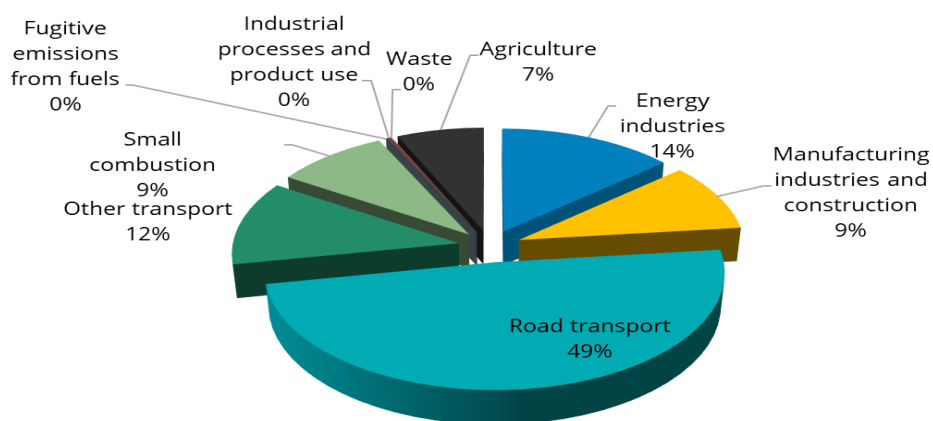


Figure 2.6.1.4 Individual sectors contribution of NO<sub>x</sub> emissions for 2017

## NM VOC Emissions

National emissions of non-methane volatile organic compounds (NMVOCs) have decreased by 53,6 % since 1990. From the year 1990 when total amount was 64,2 kt, NMVOC emissions steadily decreased to 29,8 kt in 2017. The most significant sources of NMVOC emissions in 2017 were industrial processes and product use sector (31 %) and small combustion sector (23 %). The decline in emissions since 1990 has primarily been due to reductions achieved in the road transport sector due to the introduction of vehicle catalytic converters and carbon canisters on gasoline cars for evaporative emission control, driven by tighter vehicle emission standards, combined with limits on the maximum volatility of petrol that can be sold in EU Member States, as specified in fuel quality directives. The reductions in NMVOC emissions have been enhanced by the switching from petrol to diesel cars in some EU countries, and changes in the solvent and product use subsector as a result of the introduction of legislative measures limiting the use and emissions of solvents.

Slovenia has reduced emissions since 1990 in line with its obligations under the 2001/81/EC National Emission Ceilings Directive (NECD) and Gothenburg protocol. Emissions of NMVOC were well below respective ceiling. Emissions in 2017 were 25 % below national ceiling value (40 kt NMVOC).

The 2012 revision of the Gothenburg Protocol to the UNECE LRTAP Convention and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants set emission reduction targets for NMVOC based on 2005 emission totals, to be met by countries in or before 2020. Reduction of emissions has to be 23 % compared to 2005 emissions. Emissions for Slovenia in 2017 were below a linear target path to its 2020 target by 33 % of its 2005 emission totals.

Non-methane volatile organic compounds (NMVOCs) are a collection of organic compounds that differ widely in their chemical composition but display similar behaviour in the atmosphere. NMVOCs are emitted into the atmosphere from a large number of sources including combustion activities, solvent use and production processes. Biogenic NMVOC are emitted by vegetation, with amounts dependent on the species and on temperature. NMVOCs contribute to the formation of ground-level (tropospheric) ozone, and certain species such as benzene and 1,3 butadiene are directly hazardous to human health. Quantifying the emissions of total NMVOC provides an indicator of the emissions of the most hazardous NMVOCs.

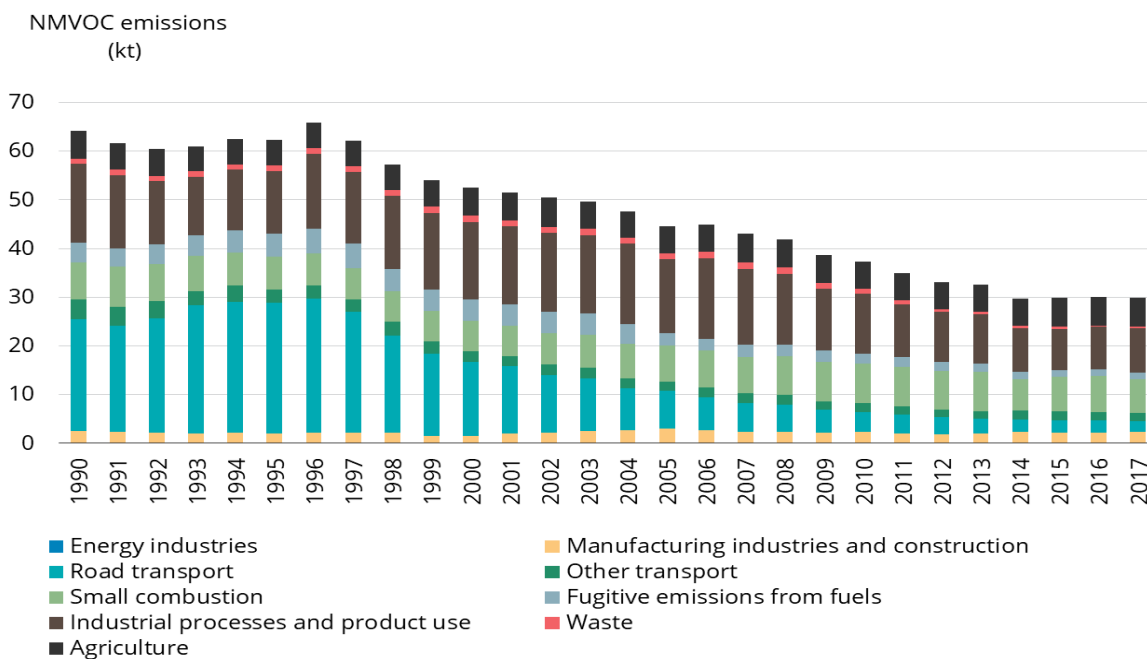


Figure 2.6.1.5 NMVOC emissions in Slovenia for the period 1990 - 2017

The main sources of NMVOC emissions in the year 2017 are industrial process and product use sector (31 %) and small combustion with a share of 23 %.

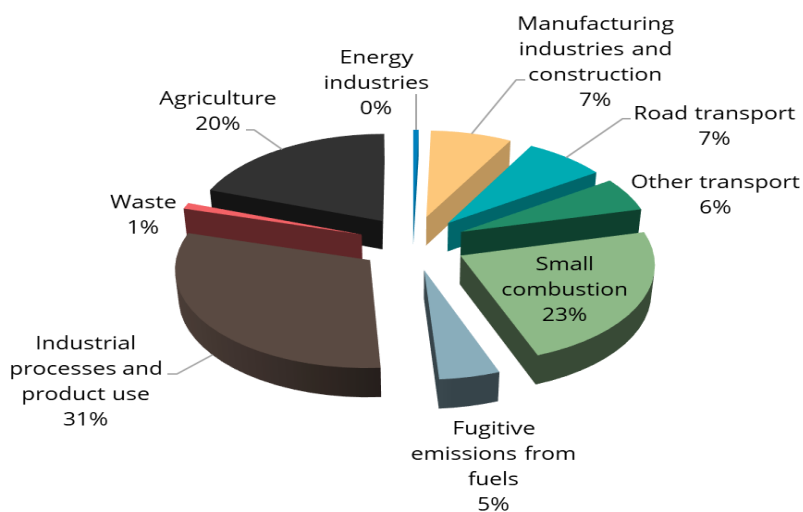


Figure 2.6.1.6 Individual sectors contribution of NMVOC emissions for 2017

## NH<sub>3</sub> Emissions

National emissions of NH<sub>3</sub> have declined by 22,4 % between the years 1986 (24,0 kt) and 2017 (18,6 kt). Agriculture was responsible for 91 % of NH<sub>3</sub> emissions in 2017. The reduction in emissions within the agricultural sector is primarily due to a reduction in livestock numbers (especially cattle), changes in the handling and management of organic manures and from the decreased use of nitrogenous fertilisers. The reductions achieved in the agricultural sector have been marginally offset by the increase in annual emissions over this period in the road-transport sector.

Total NH<sub>3</sub> emissions in 2017 were below the level of the respective 2010 ceiling (20 kt NH<sub>3</sub>). Emissions were 7 % lower than target value set in 2001/81/EC National Emission Ceilings Directive and Gothenburg protocol.

The 2012 revision of the Gothenburg Protocol to the UNECE LRTAP Convention and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants set emission reduction targets for NH<sub>3</sub> based on 2005 emission totals, to be met by countries in or before 2020. Reduction of emissions has to be 1 % compared to 2005 emissions. Emissions for Slovenia in 2017 were below a linear target path to its 2020 target by 8 % of its 2005 emission totals.

NH<sub>3</sub> contributes to acid deposition and eutrophication. The subsequent impacts of acid deposition can be significant, including adverse effects on aquatic ecosystems in rivers and lakes and damage to forests, crops and other vegetation. Eutrophication can lead to severe reductions in water quality with subsequent impacts including decreased biodiversity, changes in species composition and dominance, and toxicity effects. NH<sub>3</sub> also contributes to the formation of secondary particulate aerosols, an important air pollutant due to its adverse impacts on human health.

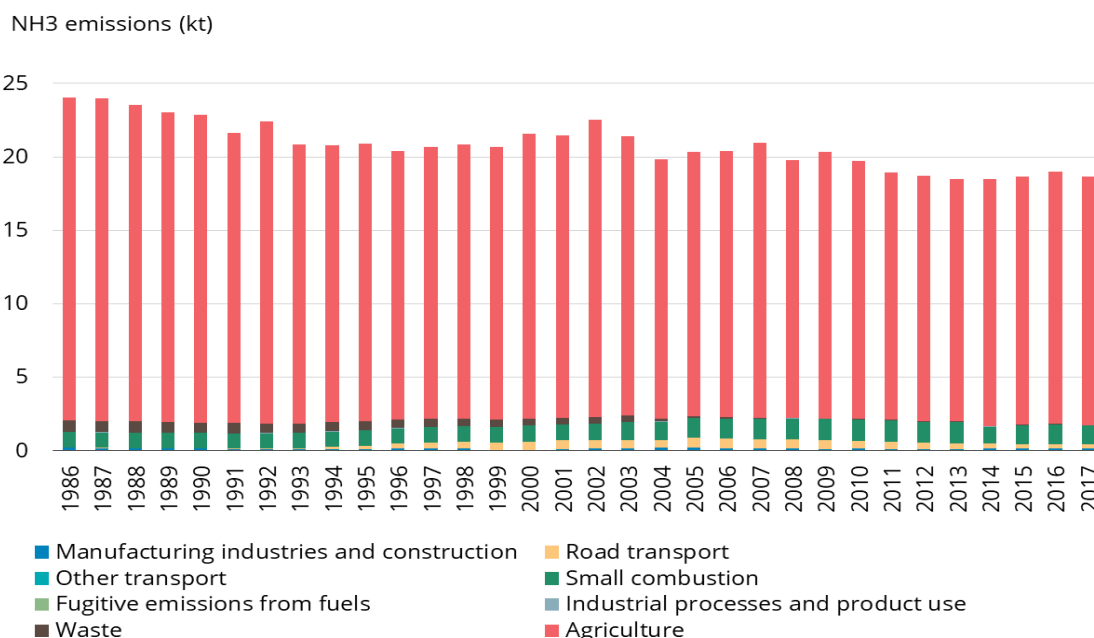
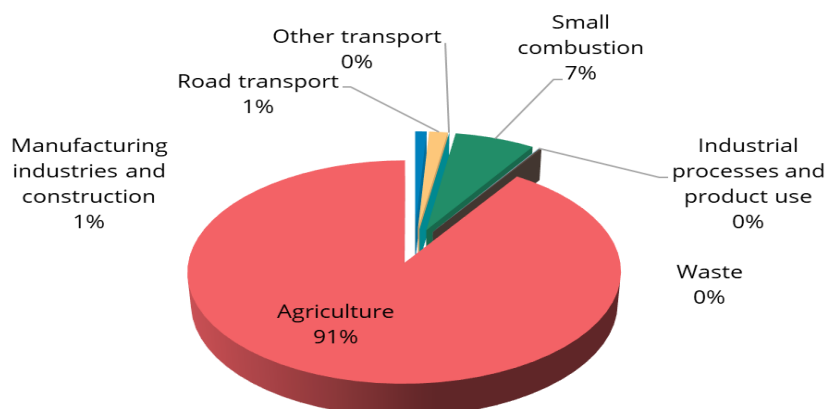


Figure 2.6.1.7 NH<sub>3</sub> emissions in Slovenia for the period 1986 – 2017



**Figure 2.6.1.8 Individual sectors contribution of NH<sub>3</sub> emissions for 2017**

## CO Emissions

National CO emissions gradually decreased from the year 1980, when total amount was 319,8 kt to 105,0 kt in 2017. Emissions were reduced by 67,2 %. This decrease has been achieved mainly as a result of the introduction of catalytic converters for gasoline vehicles, which has significantly reduced emissions of CO from the road transport sector. CO is mainly emitted from incomplete combustion. Combustion in commercial, institutional and households is responsible for the dominant share of the total CO emissions.

Emissions of carbon monoxide (as well as non-methane volatile organic compounds, nitrogen oxides and methane) contribute to the formation of ground-level (tropospheric) ozone. Ozone is a powerful oxidant and tropospheric ozone can have adverse effects on human health and ecosystems. It is a problem mainly during the summer months. High concentrations of ground-level ozone adversely affect the human respiratory system and there is evidence that long-term exposure accelerates the decline in lung function with age and may impair the development of lung function. Some people are more vulnerable to high concentrations than others, with the worst effects generally being seen in children, asthmatics and the elderly. High concentrations in the environment are harmful to crops and forests, decreasing yields, causing leaf damage and reducing disease resistance.



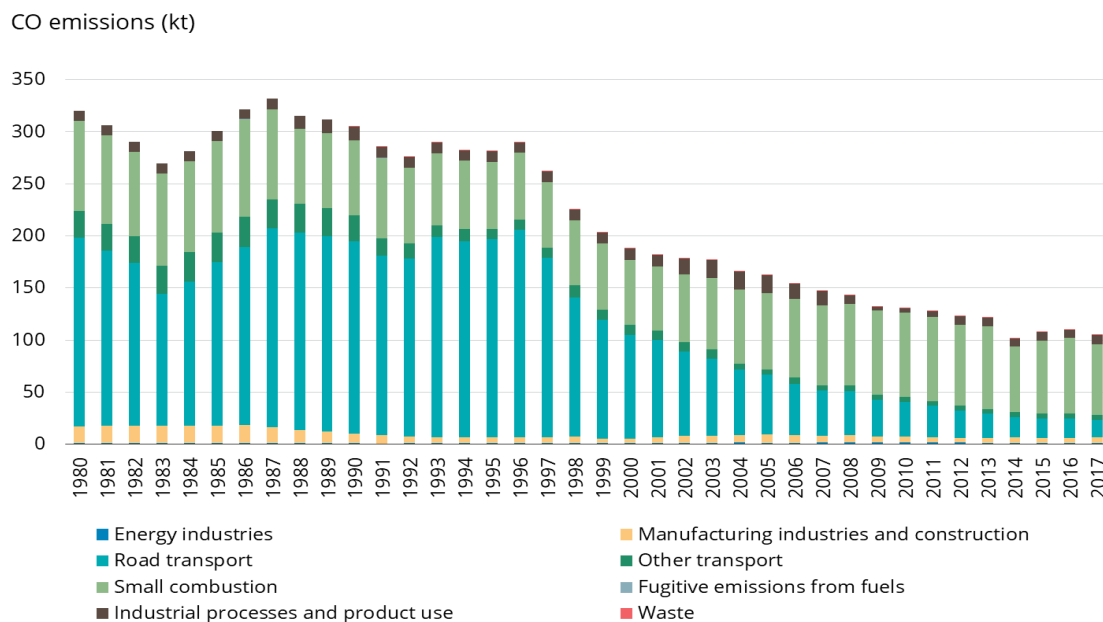


Figure 2.6.1.9 CO emissions in Slovenia for the period 1980 - 2017

In 2017, the main sources for CO emissions in Slovenia is small combustion (mainly combustion of fuel in residential sector) with a share of 64 %. Also road transport contributes significantly to the total emission of this pollutant (16 %).

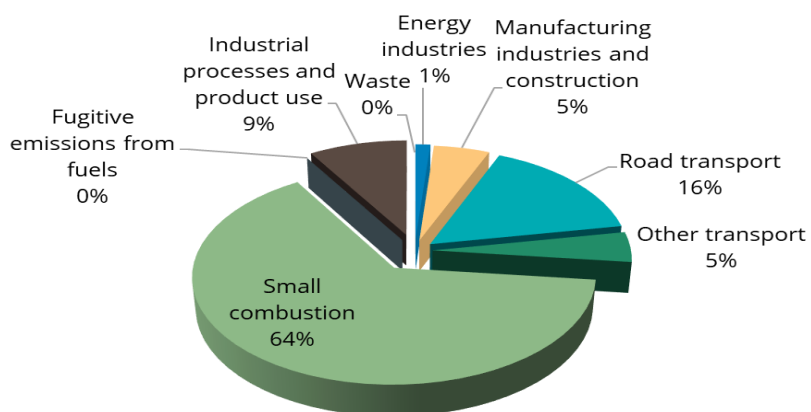


Figure 2.6.1.10 Individual sectors contribution of CO emissions for 2017

## 2.6.2 Emission Trends for Particulate Matter

The most important source of particulate matter emissions (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP and BC) has been combustion of wood in stationary residential sector. Other significant sources are road transport and use of fuel in industry production. The particulate matter emissions have increased significantly in the year 2009 due to increase of wood consumption in small combustion sector. The emission trend from year 2000 to 2017 were on the increase of PM<sub>2.5</sub> for 5,8 %, for PM<sub>10</sub> for 2,3 % and BC for 2,0 %. Emissions of TSP have decreased by 0,9 %.

The reductions in total emissions of primary PM<sub>10</sub> have not been achieved in the past decade inspite of introduction or improvement of abatement measures across the energy, road transport, and industrial sectors coupled with other developments in industrial sectors such as fuel switching from high-sulphur fuels to low-sulphur fuels, which has also contributed to decreased formation of secondary particulate matter from SO<sub>2</sub> in the atmosphere. Emissions of primary PM<sub>10</sub> are expected to decrease in the future as vehicle technologies are further improved and stationary fuel combustion emissions are controlled through abatement or use of low-sulphur fuels such as natural gas. Despite this, it is expected that within many of the urban areas across the EU, PM<sub>10</sub> concentrations will still be well above the EU air quality limit value. Substantial further reductions in emissions will therefore be needed if the limit value set in the EU's Air Quality Directive is to be reached

The 2012 revision of the Gothenburg Protocol to the UNECE LRTAP Convention and Directive (EU) 2016/2284 on the reduction of national emissions of certain atmospheric pollutants set emission reduction targets for PM<sub>2.5</sub> based on 2005 emission totals, to be met by countries in or before 2020. Reduction of emissions has to be 25 % compared to 2005 emissions. Emissions for Slovenia in 2017 were below a linear target path to its 2020 target by 14 % of its 2005 emission totals. Additional measures may therefore need to be undertaken in future years to achieve reduction target implied by the protocol.

There are no specific EU emission targets for primary PM<sub>10</sub>. However the EU National Emission Ceilings Directive (NECD) and the Gothenburg Protocol to the UNECE LRTAP Convention both set ceilings for the secondary particulate matter precursors NH<sub>3</sub>, NO<sub>x</sub> and SO<sub>x</sub> that countries must have met by 2010. NH<sub>3</sub>, NO<sub>x</sub> and SO<sub>x</sub> are ranked among secondary particulate matter precursor as well as substances which cause acidifying and eutrophication.

In recent years scientific evidence has been strengthened by many epidemiological studies that indicate there is an association between long and short-term exposure to fine particulate matter and various serious health impacts. Fine particles have adverse effects on human health and can be responsible for and/or contribute to a number of respiratory problems. Fine particles in this context refer to primary particulate matter (PM<sub>2.5</sub> and PM<sub>10</sub>) and emissions of secondary particulate matter precursors (NO<sub>x</sub>, SO<sub>x</sub> and NH<sub>3</sub>). Primary PM<sub>2.5</sub> and PM<sub>10</sub> refers to fine particles (defined as having diameter of 2.5 µm or 10 µm or less, respectively) emitted directly to the atmosphere. Secondary particulate matter precursors are pollutants that are partly transformed into particles by photo-chemical reactions in the atmosphere. A large fraction of the urban population is exposed to levels of fine particulate matter in excess of limit values set for the protection of human health. There have been a number of recent policy initiatives that aim to control particulate concentrations and thus protect human health.

**Table 2.6.2.1 National total emissions and emission trends for the period 2000-2017 for particulate matter**

Year	Emissions (kt)			
	PM <sub>2.5</sub>	PM <sub>10</sub>	TSP	BC
2000	10,9	12,7	15,3	2,0
2001	11,1	13,0	15,6	2,1
2002	11,7	13,7	16,3	2,2
2003	12,2	14,1	16,7	2,3
2004	12,8	14,6	17,2	2,5
2005	13,3	15,3	18,3	2,6
2006	13,2	15,1	18,1	2,6
2007	13,2	15,1	18,0	2,6
2008	14,0	16,0	18,9	2,8
2009	13,4	15,1	17,5	2,6
2010	13,4	15,1	17,6	2,6
2011	13,3	15,0	17,5	2,5
2012	12,8	14,5	16,9	2,4
2013	12,9	14,5	16,7	2,4
2014	11,0	12,4	14,5	2,1
2015	11,6	13,1	15,2	2,1
2016	11,9	13,4	15,6	2,2
2017	11,5	13,0	15,1	2,1
<b>Trend (%)</b>	<b>+5,8 %</b>	<b>+2,3 %</b>	<b>-0,9 %</b>	<b>+2,0 %</b>

### PM<sub>10</sub> Emissions

In the year 2017 the total amount of primary PM<sub>10</sub> (sub-10µm particulate matter) emissions accounted to 13,0 kt. Emissions in the year 2000 were 12,7 kt. The most important source of primary PM<sub>10</sub> emissions in 2017 was small combustion sector which includes combustion-related emissions from sources such as heating of residential and commercial properties mainly wood consumption in residential sector (66 %). Other important sectors are road transport (10 %) and fuel used in manufacturing industries and construction (8 %).

Emissions of primary PM<sub>10</sub> have increased from 2000 to 2017 by 2 %. Increase of emissions was the most pronounced in small combustion sector and in road transport sector. Bigger fuel consumption in recent years is the reason for increase of particle emissions, in spite of improvements in vehicle technologies. Increase of emissions in 2009 in residential sector is due to biomass burning in inefficient stoves. The use of biomass in households increased due to favourable price of biomass compared to other fuels as well as state measures to promote renewable energy sources. The decrease in emissions in the past two years was due to significantly reduced emissions from residential combustion. Warmer winter and improved thermal insulation of buildings contributed to lower fuel consumption.

Other factors which contributed to the reduction of primary PM<sub>10</sub> emissions in some sectors are: improvements in the performance of particulate abatement equipment at industrial combustion facilities (coal-fired power stations), a fuel shift from the use of coal in the energy industries, industrial and domestic sectors to cleaner burning fuels such as gas, cleaner stoves for domestic heating, introduction of particle filters on new vehicles (driven by the legislative EURO standards).

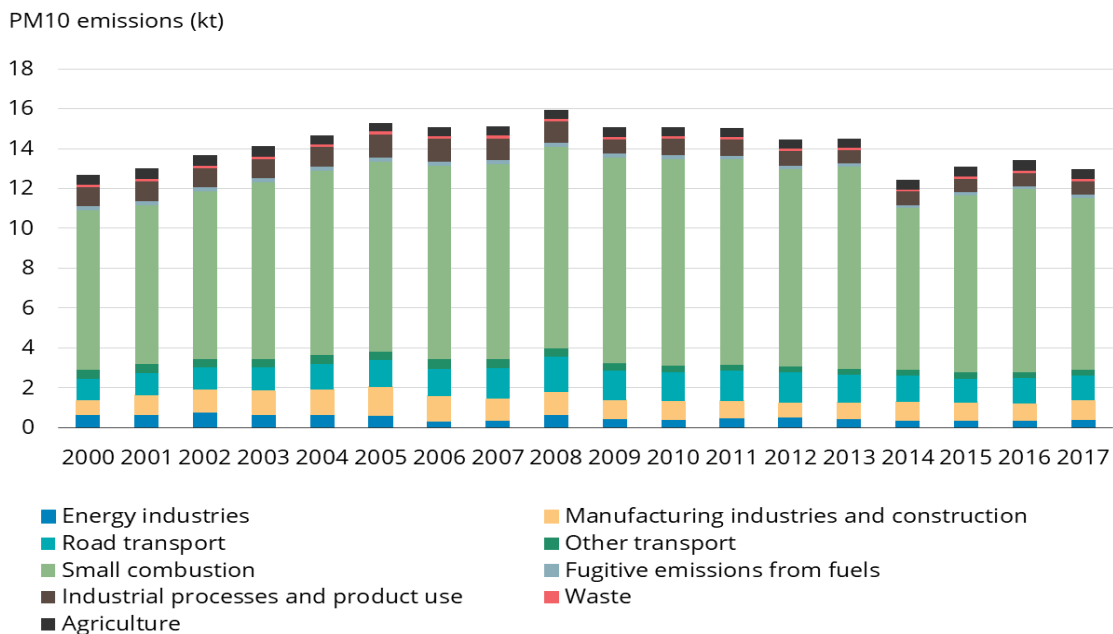


Figure 2.6.2.1 PM<sub>10</sub> emissions in Slovenia for the period 2000 - 2017

The main source for PM<sub>10</sub> emissions in the year 2017 was small combustion sector mainly wood consumption in residential sector with a share of about 66 %, followed by road transport with 10 %.

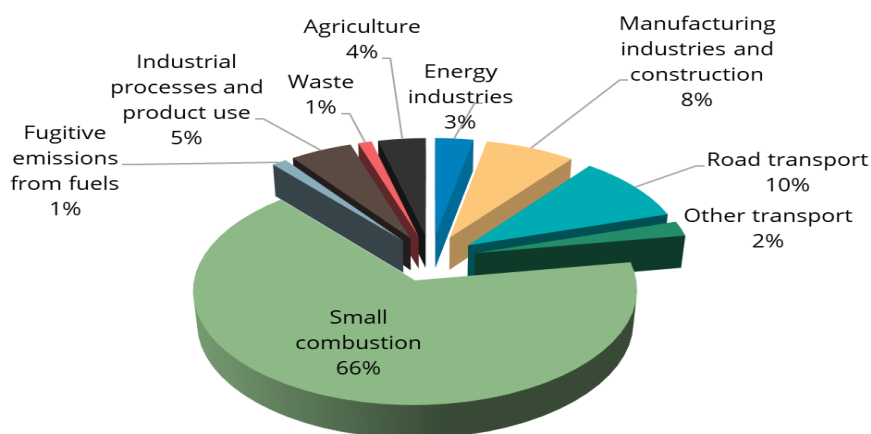


Figure 2.6.2.2 Individual sectors contribution of PM<sub>10</sub> emissions for 2017

## PM<sub>2.5</sub> Emissions

National PM<sub>2.5</sub> emissions increased by 6 % from the year 2000, when total amount was 10,9 kt, to 11,5 kt in 2017.

The PM<sub>2.5</sub> emissions have increased in 2009 in stationary residential sector due to increase of wood consumption. Increasing consumption of biomass is probably a result of economic crisis and a high price of petroleum products as well as state measures to promote renewable energy sources. The decrease in emissions in 2014 and 2015 was due to significantly reduced emissions from residential combustion. Warmer winter and improved thermal insulation of buildings contributed to lower fuel consumption.

Far most important source of PM<sub>2.5</sub> emissions in the year 2017 was small combustion sector with a share of 73 %, followed by road transport and fuel used in manufacturing industries and construction with 8 %.

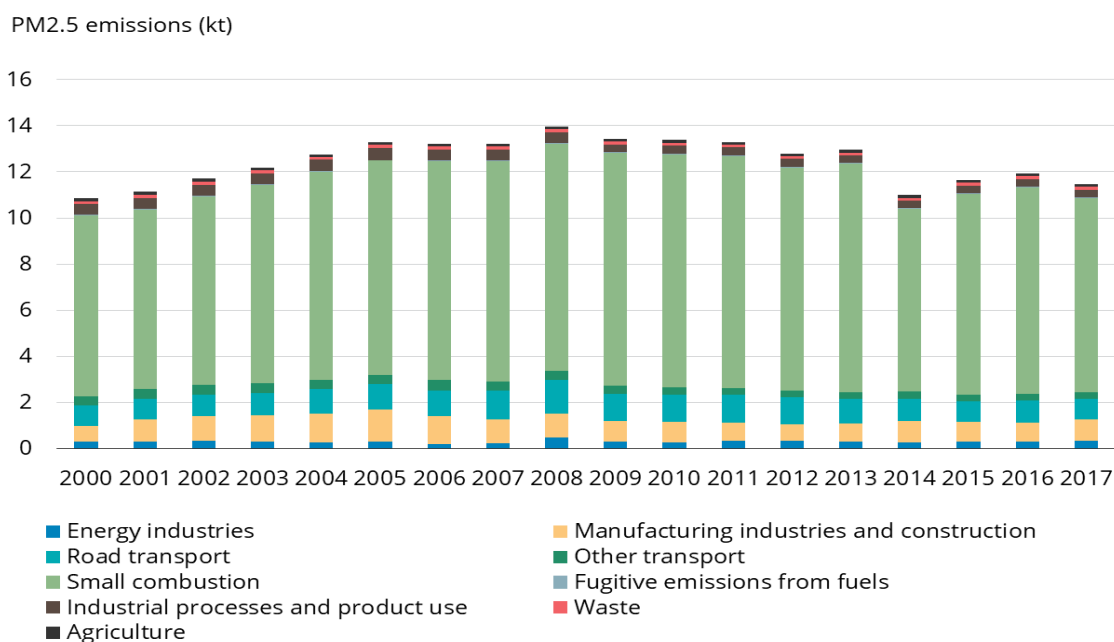
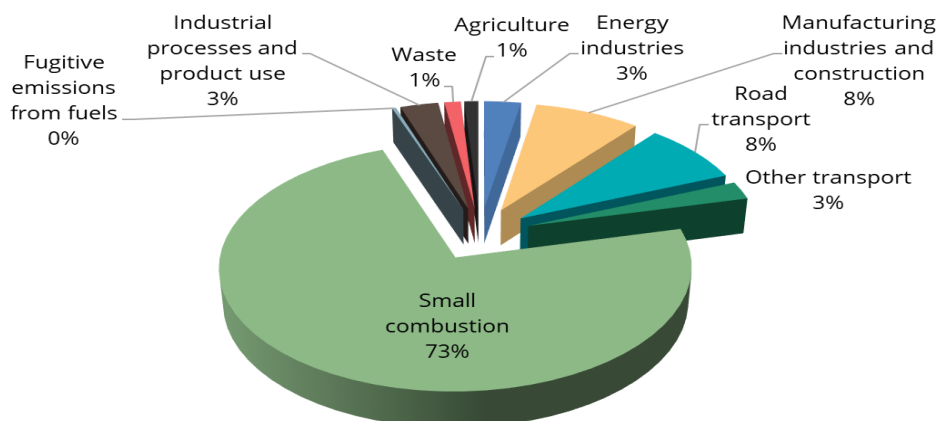


Figure 2.6.2.3 PM<sub>2.5</sub> emissions in Slovenia for the period 2000 – 2017



**Figure 2.6.2.4 Individual sectors contribution of PM<sub>2.5</sub> emissions for 2017**

### TSP Emissions

National total suspended particulate (TSP) emissions have decreased from the year 2000, when total amount was 15,3 kt to 15,1 kt in 2017. Emissions were decreased by 1 % mainly due to decrease of emissions in production of energy. The TSP emissions have increased in 2009 in stationary residential sector due to increase of wood consumption. Increasing consumption of biomass is probably a result of economic crisis and a high price of petroleum products as well as state measures to promote renewable energy sources. The decrease in emissions in 2014 and 2015 was due to significantly reduced emissions from residential combustion. Warmer winter and improved thermal insulation of buildings contributed to lower fuel consumption.

The main source of TSP emissions in the year 2017 was small combustion sector with a share of 59 %. Contribution of road transport was 11 %.

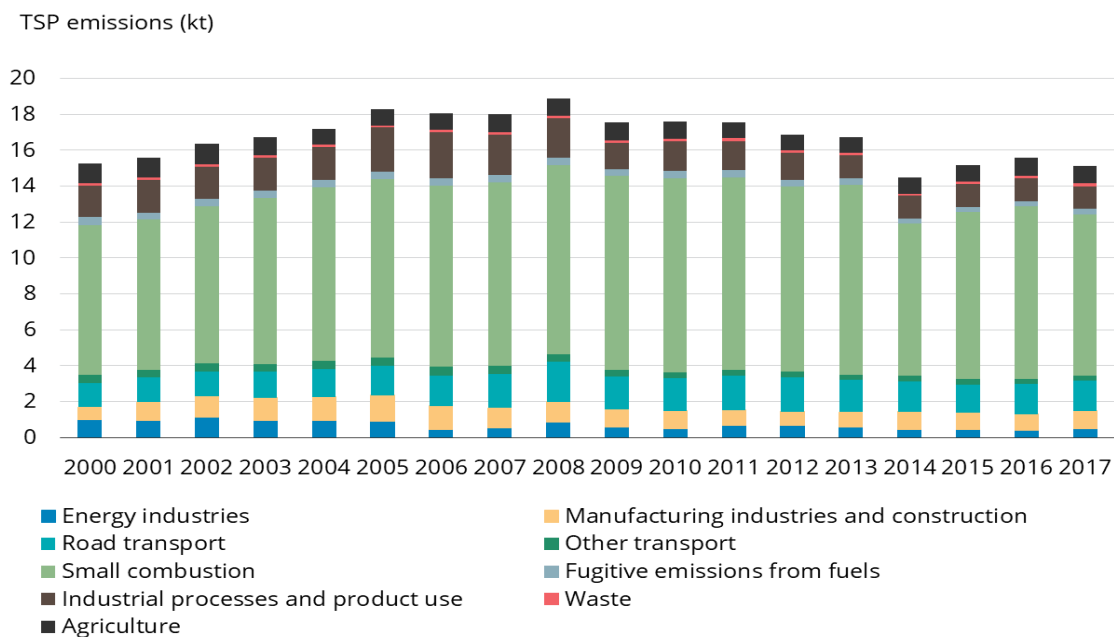


Figure 2.6.2.5 TSP emissions in Slovenia for the period 2000 – 2017

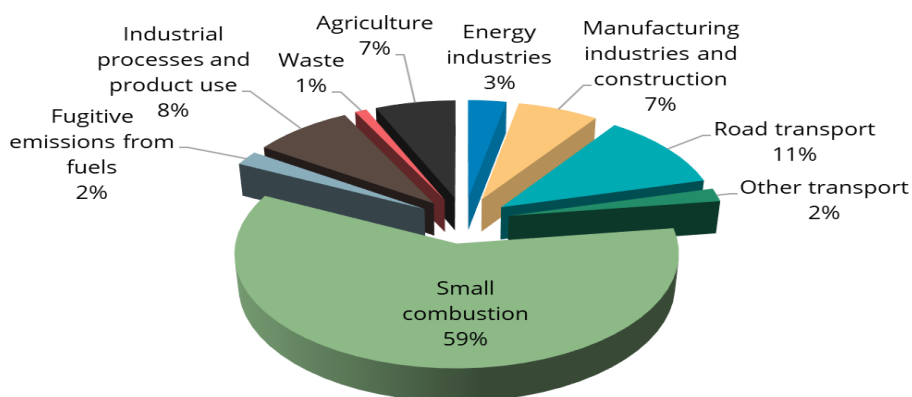


Figure 2.6.2.6 Individual sectors contribution of TSP emissions for 2017

### Black carbon Emissions

National black carbon (BC) emissions increased from the year 2000, when total amount was 2,0 kt to 2,1 kt in 2017. Emissions were increased by 2 % mainly in energy industries. Far most important source of BC emissions in the year 2017 was small combustion sector with a share of 61 %, followed by road transport with 20 %, fuel consumption in manufacturing and construction (11 %) and other transport (8 %).

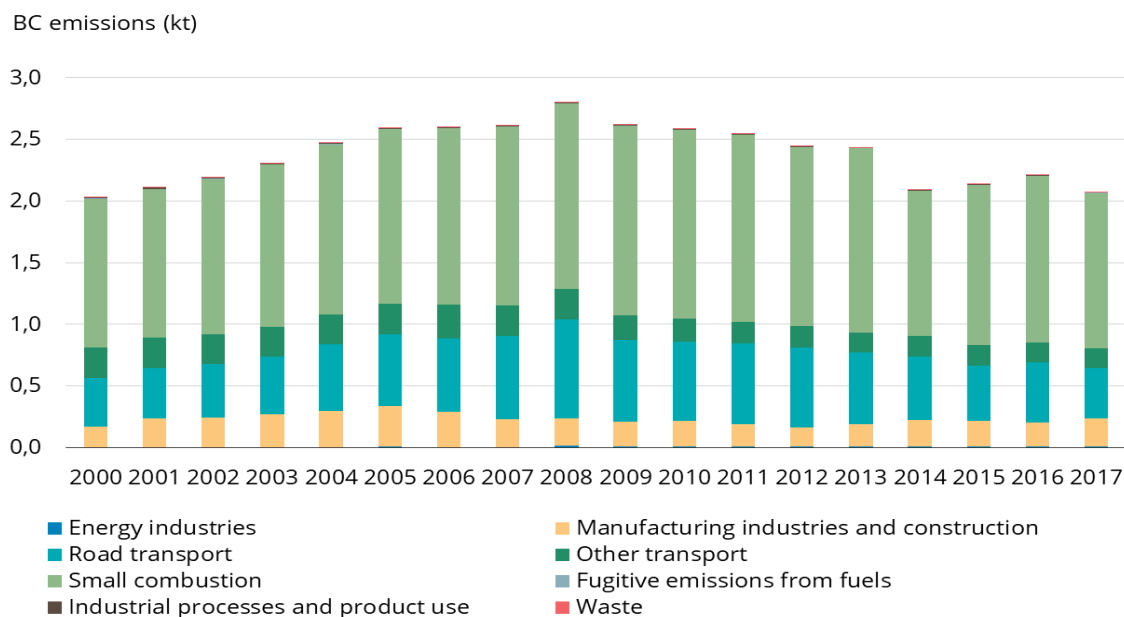


Figure 2.6.2.7 BC emissions in Slovenia for the period 2000 – 2017

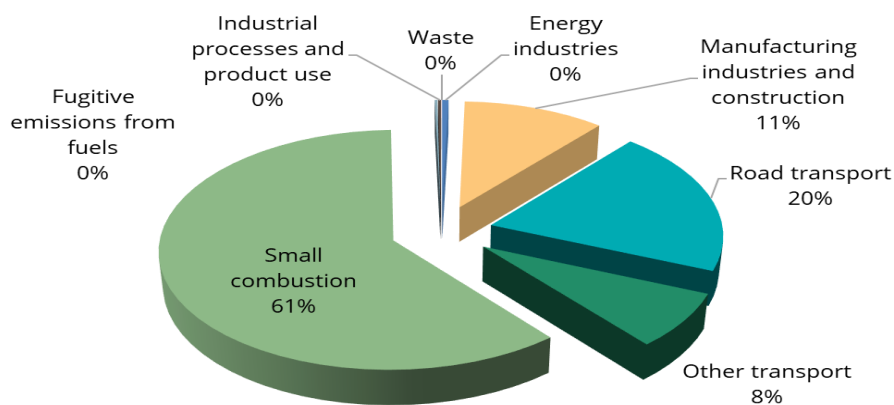


Figure 2.6.2.8 Individual sectors contribution of BC emissions for 2017



### 2.6.3 Emission Trends for Heavy Metals

In general, the most important sources of heavy metals (Pb, Cd and Hg) emissions have been production processes, combustion of fossil fuels and road transport. Emissions of lead have decreased by 98 %, mercury by 50 % and cadmium by 15 % between 1990 and 2017. The most significant sources of heavy metals are from industrial facilities and energy-related fuel combustion. The reason for the reduced emissions is mainly increased use of gas cleaning devices at power plants. Lead and cadmium emissions have also both decreased from certain industrial processes, such as metal refining and smelting activities, reflecting improved pollution abatement control and also as a result of economic restructuring and the closure of older and more polluting industrial facilities. In the case of mercury, the observed decrease in emissions may be largely attributed to improved controls on mercury in industrial processes (installation of pollution control equipment – flue gas desulphurization system and the decline of coal use as a result of fuel switching. The promotion of unleaded petrol has been the main reason for decline of Pb emissions. Leaded petrol was phased out in Slovenia in the year 2002. Nevertheless, the road transport sector still remains a principal source of lead, contributing around 46 % of total lead emissions. However since 2002 little progress has been made in reducing emissions further. 98% of the total reduction from 1990 emissions of lead had been achieved by 2002. Residual lead in fuel, from engine lubricants and parts, and from tyre and brake wear contribute to the on-going lead emissions from this sector.

Heavy metals such as cadmium, lead and mercury are recognised as being toxic to biota. All are prone to biomagnification, being progressively accumulated higher up the food chain, such that bioaccumulation in lower organisms at relatively low concentrations can expose higher consumer organisms, including humans, to potentially harmful concentrations. In humans they are also of direct concern because of their toxicity, their potential to cause cancer and their potential ability to cause harmful effects at low concentrations. The relative toxic/carcinogenic potencies of heavy metals are compound specific, but exposure to heavy metals has been linked with developmental retardation, various cancers and kidney damage. Metals are persistent throughout the environment. These substances tend not just to be confined to a given geographical region, and thus are not always open to effective local control. For example, in the case of cadmium, much is found in fine particles which do not readily dry-deposit, and therefore have long residence times in the atmosphere and are subject to long-range transport processes.

Slovenia in 2017 did not exceed emission levels set in protocol on heavy metals. Emissions are much below values from the reference year 1990.

Emissions of additional heavy metals (As, Cr, Cu, Ni, Se, Zn) have been estimated for the first time in this annual submission. Emissions of As, Cr, Ni, Se have decreased by 27, 11, 48 and 32 %, respectively, from the year 1990 to 2017. Emissions of Cu and Zn have increased by 68 % and 17 % between 1990 and 2017.

**Table 2.6.3.1 National total emissions and emission trends for the period 1990 - 2017 for Pb, Cd and Hg**

Year	Emissions (t)		
	Pb	Cd	Hg
1990	342,2	0,70	0,33
1991	300,3	0,59	0,30
1992	292,2	0,61	0,30
1993	308,1	0,55	0,28
1994	308,0	0,54	0,27

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1995	192,4	0,54	0,22
1996	76,5	0,52	0,20
1997	66,2	0,57	0,22
1998	50,9	0,60	0,23
1999	43,4	0,56	0,21
2000	36,3	0,60	0,20
2001	20,4	0,63	0,19
2002	9,5	0,66	0,22
2003	9,6	0,68	0,21
2004	9,2	0,70	0,19
2005	9,6	0,72	0,19
2006	9,5	0,73	0,18
2007	9,4	0,76	0,18
2008	10,0	0,77	0,18
2009	8,2	0,67	0,16
2010	8,9	0,73	0,19
2011	8,9	0,69	0,19
2012	8,5	0,61	0,18
2013	7,8	0,62	0,18
2014	7,1	0,55	0,16
2015	7,3	0,58	0,16
2016	7,3	0,60	0,16
2017	7,2	0,60	0,16
<b>Reduction trend (%)</b>	<b>-97,9 %</b>	<b>-15,0 %</b>	<b>-49,7 %</b>

Table 2.6.3.2 National total emissions and emission trends for the period 1990 - 2017 for additional heavy metals: As, Cr, Cu, Ni, Se, Zn

Year	Emissions (t)					
	As	Cr	Cu	Ni	Se	Zn
1990	0,93	1,56	3,54	2,82	2,92	18,5
1991	0,83	1,42	3,27	2,55	2,79	17,2
1992	0,91	1,45	3,29	1,99	2,97	16,7
1993	0,85	1,40	3,44	2,38	2,76	16,5
1994	0,82	1,35	3,67	2,21	2,58	16,8
1995	0,81	1,36	3,97	2,26	2,50	16,9
1996	0,74	1,38	4,45	2,73	2,27	17,2
1997	0,81	1,46	4,63	2,80	2,42	17,3
1998	0,86	1,51	4,24	2,78	2,54	17,0
1999	0,77	1,40	4,03	2,64	2,28	15,4
2000	0,81	1,44	4,22	2,61	2,35	15,7
2001	0,87	1,52	4,38	2,65	2,54	16,8
2002	0,92	1,60	4,47	2,62	2,66	17,7
2003	0,87	1,60	4,63	2,55	2,51	18,9
2004	0,88	1,64	4,76	2,49	2,55	19,8
2005	0,88	1,68	5,18	2,48	2,54	21,1
2006	0,91	1,69	5,29	2,37	2,61	21,2
2007	0,94	1,74	5,80	2,09	2,68	21,4

<b>2008</b>	0,92	1,83	6,80	2,53	2,59	23,2
<b>2009</b>	0,85	1,65	5,63	2,19	2,45	21,3
<b>2010</b>	0,87	1,69	5,85	2,23	2,50	22,3
<b>2011</b>	0,87	1,64	6,01	2,08	2,54	22,6
<b>2012</b>	0,80	1,50	6,05	1,77	2,42	22,0
<b>2013</b>	0,78	1,49	5,62	1,75	2,33	22,2
<b>2014</b>	0,60	1,27	5,42	1,43	1,77	20,1
<b>2015</b>	0,63	1,34	5,48	1,44	1,87	21,1
<b>2016</b>	0,68	1,41	5,76	1,56	2,02	21,8
<b>2017</b>	0,67	1,40	5,95	1,46	1,99	21,7
<b>Trend (%)</b>	<b>-27,1 %</b>	<b>-10,5 %</b>	<b>+68,2 %</b>	<b>-48,0 %</b>	<b>-31,7 %</b>	<b>+17,0 %</b>

## Lead Emissions

National lead (Pb) emissions decreased from the year 1990, when total amount was 342,2 t to 7,2 t in 2017. Emissions of lead have declined by 98,0 % between 1990 and 2017, primarily due to reductions made in emissions from the road transport sector. The promotion of unleaded petrol was the main reason for huge reduction. The leaded petrol was phased out in Slovenia in July 2002. The large reduction of lead emissions from the road transport sector (of nearly 99 %) has been responsible for the vast majority of the overall reduction of lead emissions since 1990. Nevertheless, the road transport sector still remains an important source of lead, contributing 48 % to total national lead emission. Pb emissions decreased in 1995 and 1996 due to lowering levels of lead content in gasoline. Residual lead in fuel, from engine lubricants and parts, and from tyre and brake wear contribute to the on-going lead emissions from this sector.

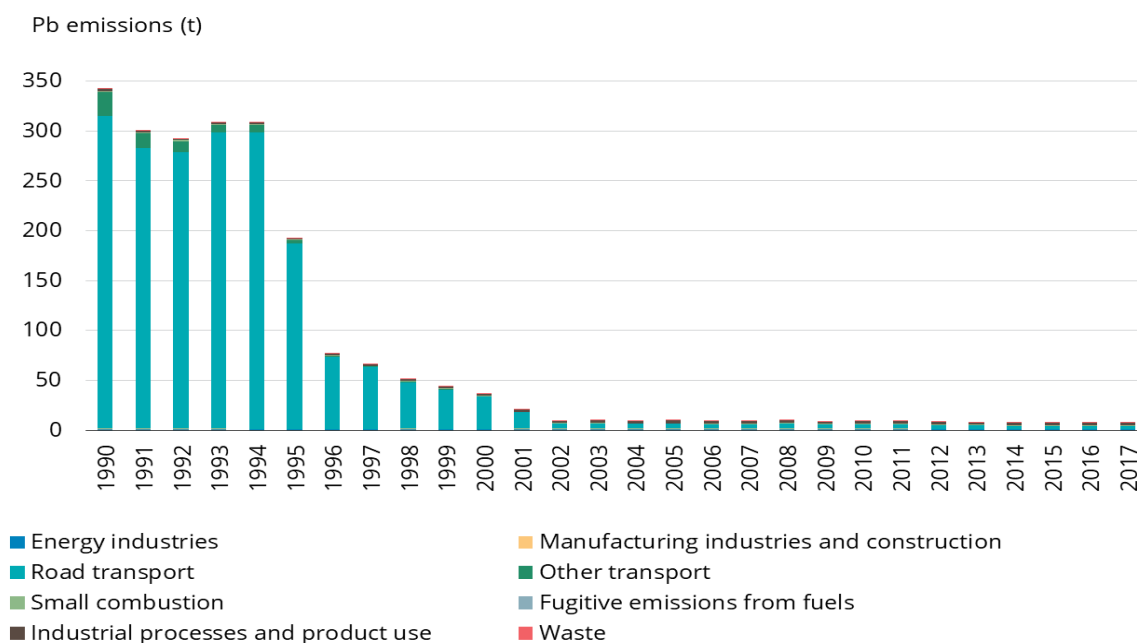


Figure 2.6.3.1 Pb emissions in Slovenia for the period 1990 – 2017

The main source for Pb emissions in the year 2017 was road transport with a share of 46 %. Contribution of industrial processes sector was 31 %.

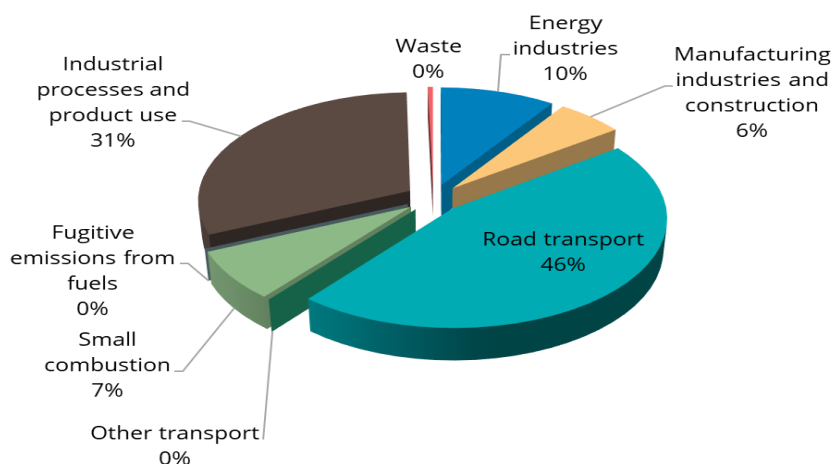


Figure 2.6.3.2 Individual sectors contribution of Pb emissions for 2017

### Cadmium Emissions

National cadmium (Cd) emissions decreased from the year 1990, when total amount was 0,70 t to 0,60 t in 2017. Emissions were reduced between 1990 and 2017 by 15 %. Decline in emissions is largely due to improvements in abatement technologies for wastewater treatment, incinerators and in metal refining and smelting facilities, coupled with the effect of European commission directives and regulations mandating reductions and limits on heavy metal emissions (e.g. the IED, IPPC directive and associated permitting conditions). The main source of Cd emissions in the year 2017 was small combustion sector with a share of 42 %. Contribution of industrial processes was 29 %.

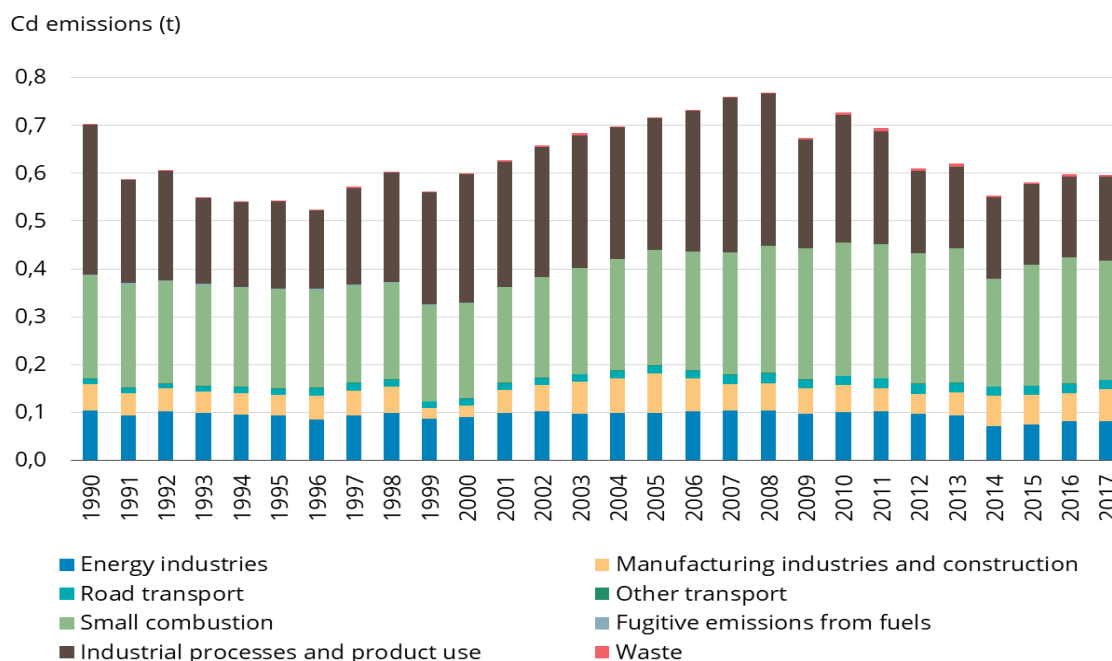


Figure 2.6.3.3 Cd emissions in Slovenia for the period 1990 –2017

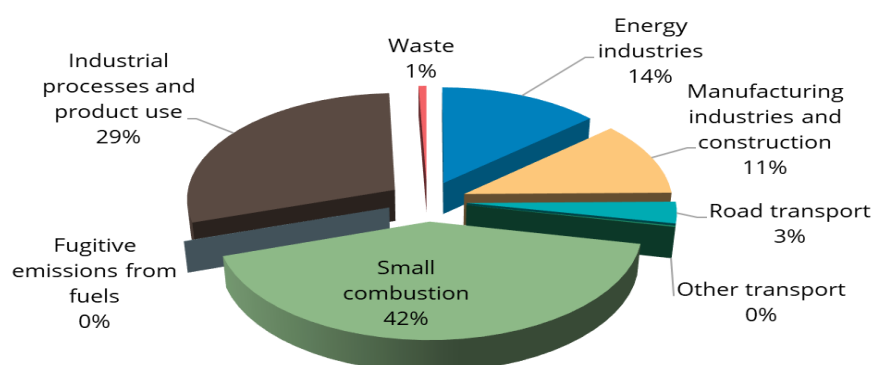


Figure 2.6.3.4 Individual sectors contribution of Cd emissions for 2017

### Mercury Emissions

National emissions of mercury (Hg) decreased from 0,33 t in year 1990 to 0,16 t in 2017. Emissions of mercury have declined by 50 % between 1990 and 2017. Since 1990 the largest reduction in mercury emissions has been achieved by the energy production and distribution sector in public power and heat generation. Mercury emissions from this sector are closely linked to the use of coal, which contains mercury as a contaminant. Past changes in fuel use within this sector since 1990, particularly fuel switching in many countries from coal to gas and other energy sources, closure of older inefficient coal-burning plants, and improved pollution

abatement equipment are mainly responsible for the past decreases in emissions from this sector.

The main source of Hg emissions in the year 2017 was industrial processes with a share of 30 %, followed by waste sector with a share of 23 %. Production of public electricity and heat contributes about 21 % to total Hg emissions.

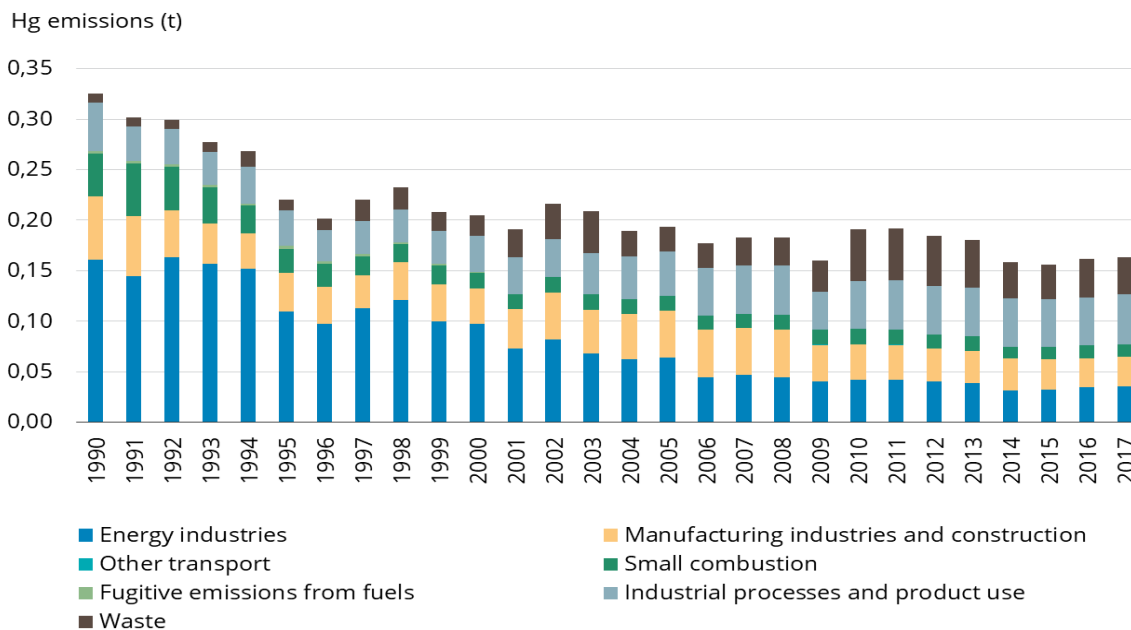


Figure 2.6.3.5 Hg emissions in Slovenia for the period 1990 – 2017

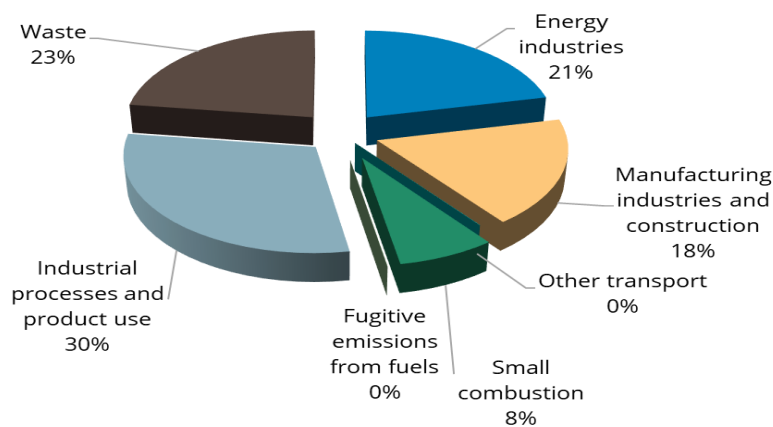


Figure 2.6.3.6 Individual sectors contribution of Hg emissions for 2017

### Arsenic Emissions

National emissions of arsenic (As) decreased from 0,93 t in year 1990 to 0,67 t in 2017. Emissions of arsenic have declined by 27 % between 1990 and 2017. Significant drop of emissions in 2014 was due to smaller use of fuels in energy and small combustion sector. The main source of As emissions in the year 2017 was production of public electricity and heat with a share of 93 %.

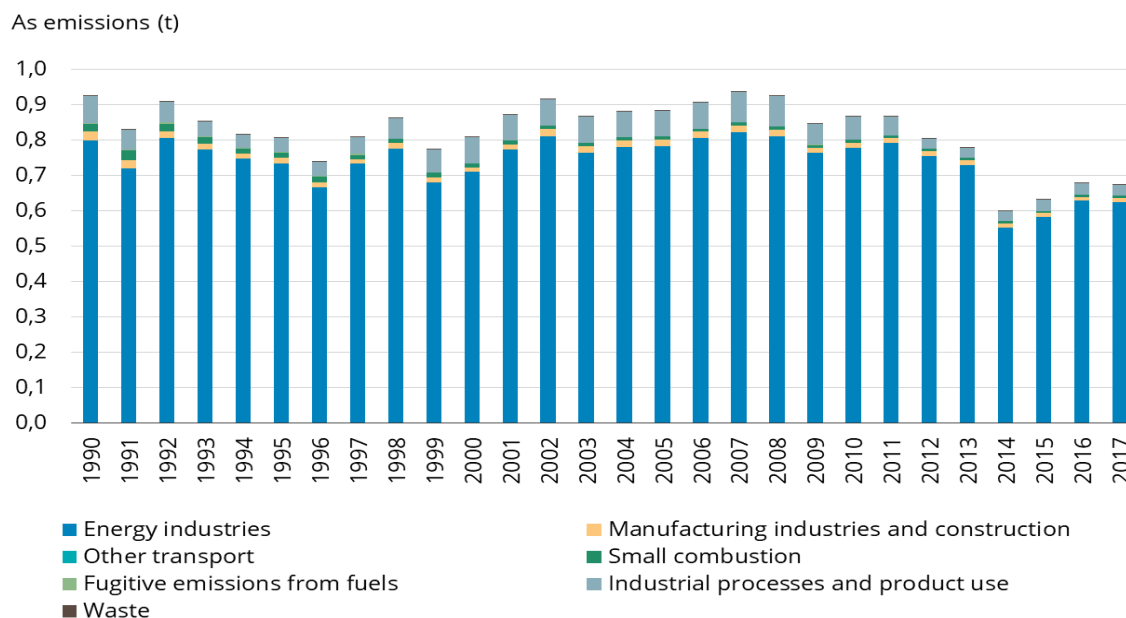


Figure 2.6.3.7 As emissions in Slovenia for the period 1990 – 2017

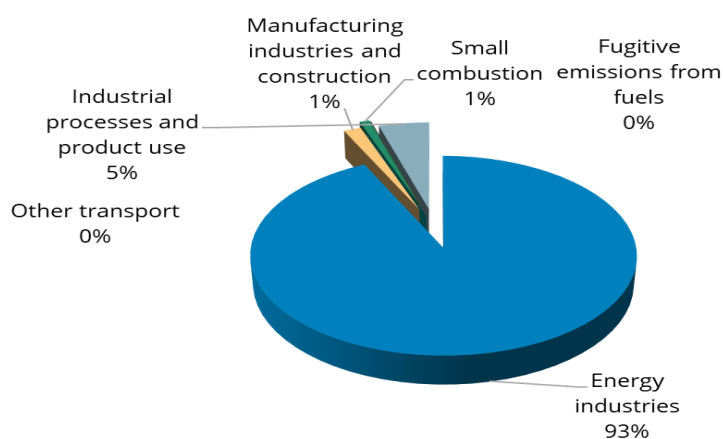


Figure 2.6.3.8 Individual sectors contribution of As emissions for 2017

### Chromium Emissions

National chromium (Cr) emissions decreased from the year 1990, when total amount was 1,56 t to 1,40 t in 2017. Emissions were reduced between 1990 and 2017 by 11 %. Significant drop of emissions in 2014 was due to smaller use of fuels in energy and small combustion sector. The main source of Cr emissions in the year 2017 was small combustion with a share of 34 %, followed by energy industries with 29 % and road transport with 20 %.

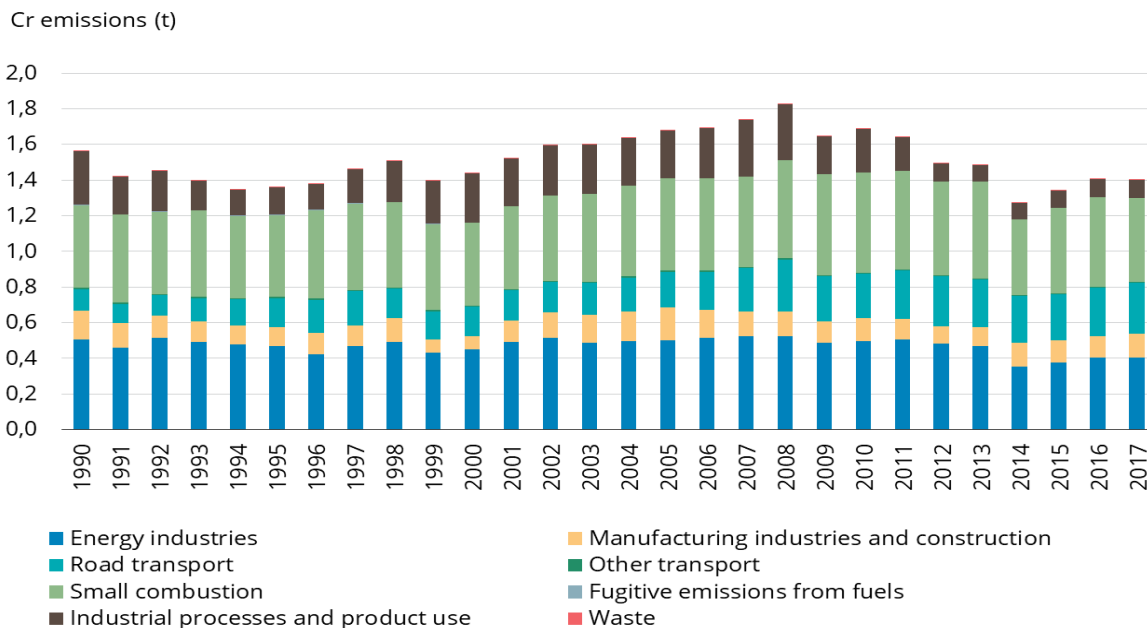


Figure 2.6.3.9 Cr emissions in Slovenia for the period 1990 –2017

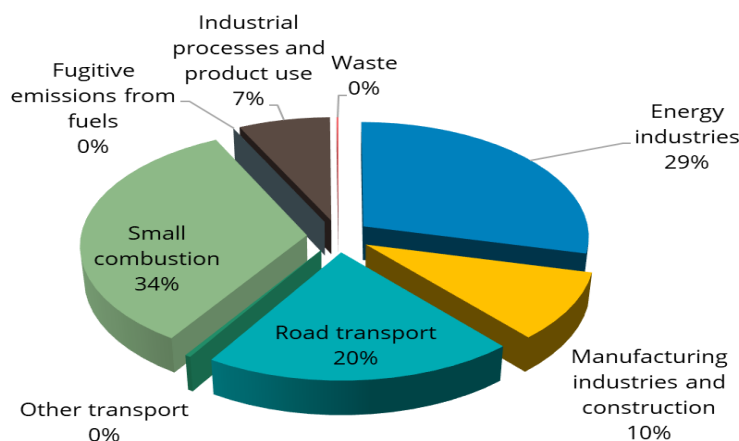


Figure 2.6.3.10 Individual sectors contribution of Cr emissions for 2017



### Copper Emissions

National copper (Cu) emissions increased from the year 1990, when total amount was 3,54 t to 5,95 t in 2017. Emissions were increased between 1990 and 2017 by 68 %. The most important source of Cu emissions in the year 2017 was road transport with a share of 88 %.

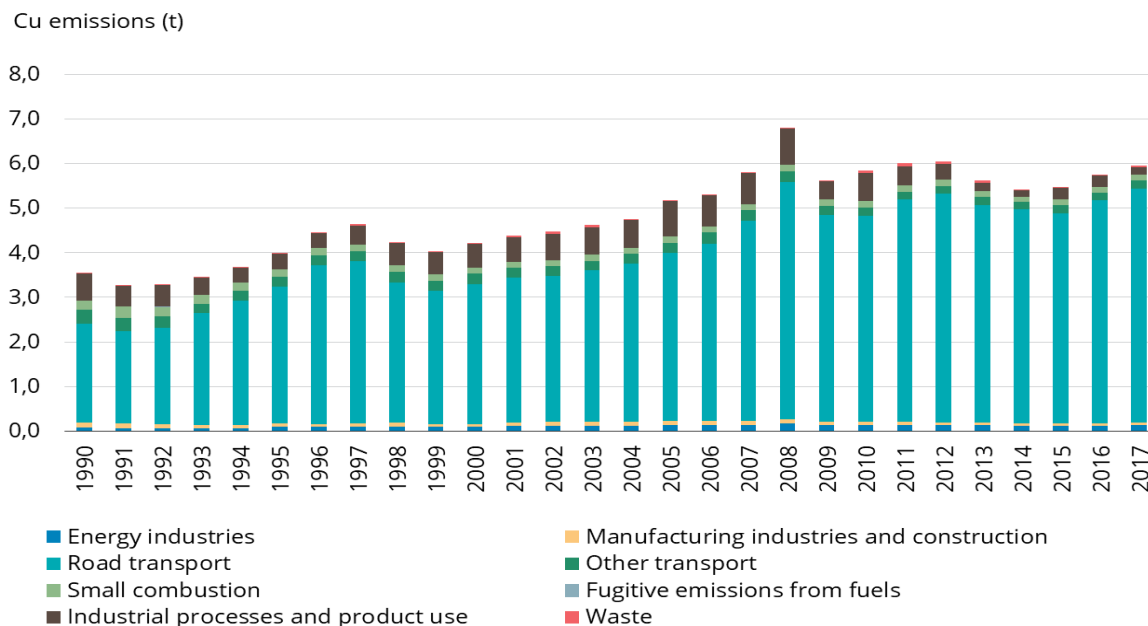


Figure 2.6.3.11 Cu emissions in Slovenia for the period 1990 –2017

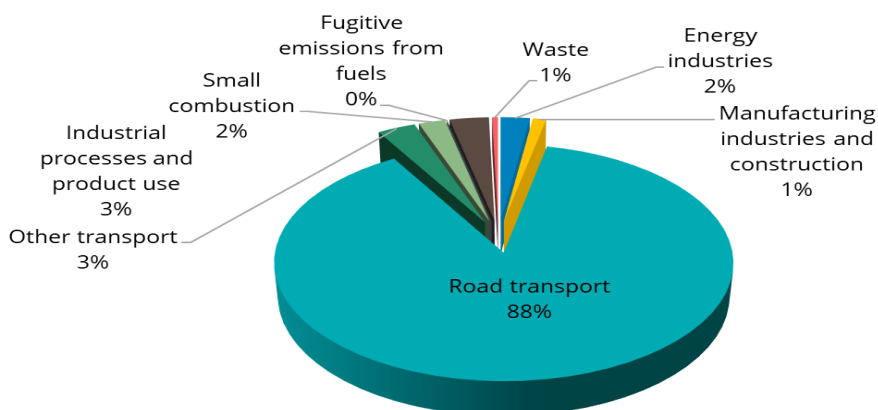


Figure 2.6.3.12 Individual sectors contribution of Cu emissions for 2017

### Nickel Emissions

National nickel (Ni) emissions decreased from the year 1990, when total amount was 2,82 t to 1,46 t in 2017. Emissions were decreased between 1990 and 2017 by 48 %. Significant drop of emissions in 2014 was due to smaller use of fuels in energy and small combustion sector. The main source of Ni emissions in the year 2017 was industrial processes and product use sector with a share of 37 %, followed by energy industries with 30 % and small combustion with 26 %.

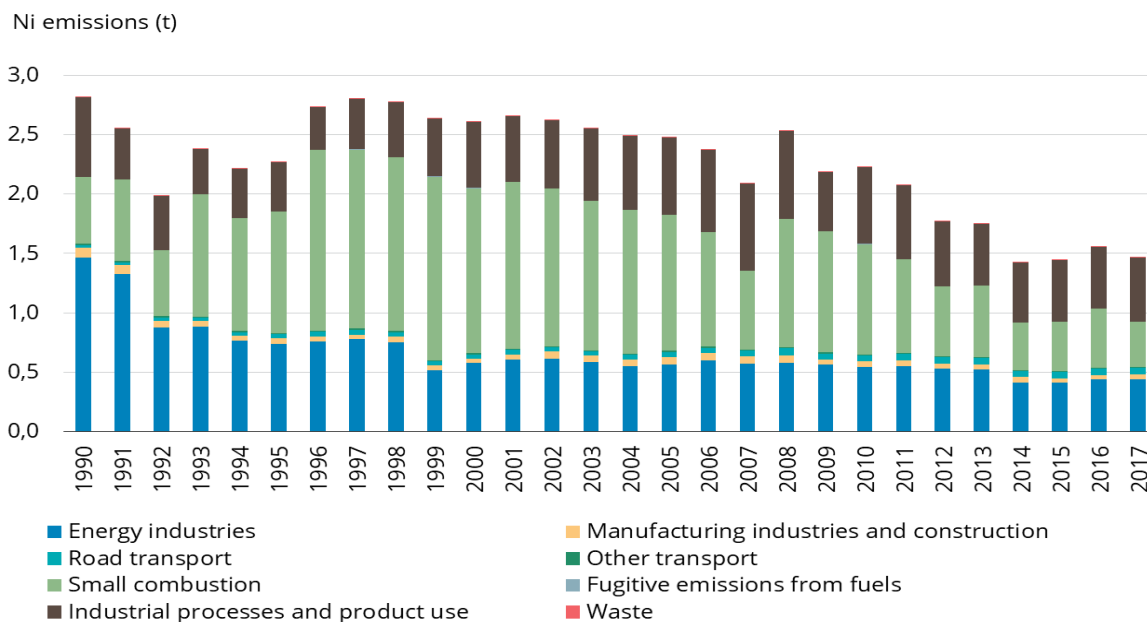


Figure 2.6.3.13 Ni emissions in Slovenia for the period 1990 –2017

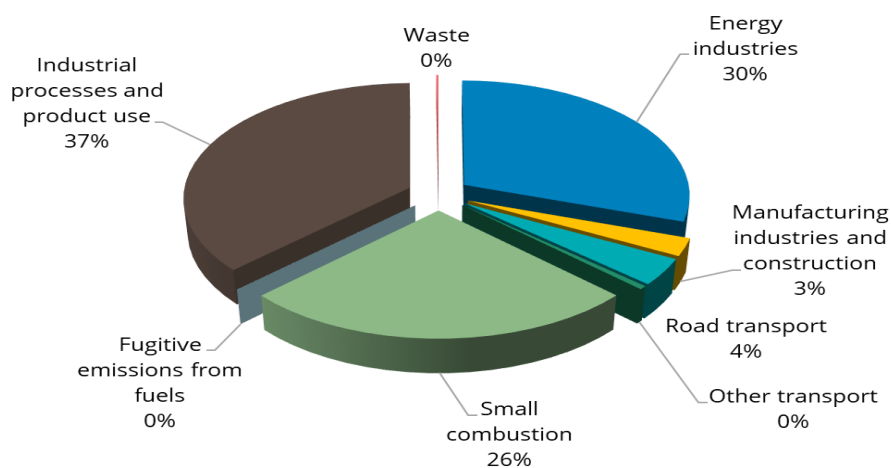


Figure 2.6.3.14 Individual sectors contribution of Ni emissions for 2017

### Selenium Emissions

National selenium (Se) emissions decreased from the year 1990, when total amount was 2,92 t to 1,99 t in 2017. Emissions were decreased between 1990 and 2017 by 32 %. Significant drop of emissions in 2014 was due to smaller use of fuels in energy and small combustion sector. The most important source of Se emissions in the year 2017 was production of energy and heat with a share of 96 %.

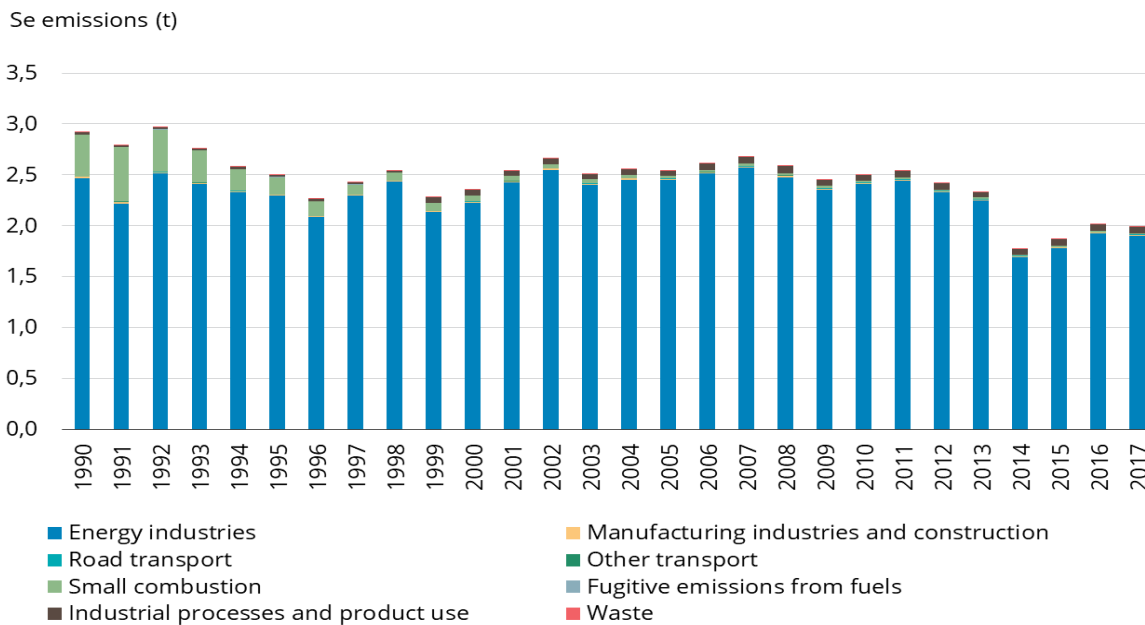


Figure 2.6.3.15 Se emissions in Slovenia for the period 1990 –2017

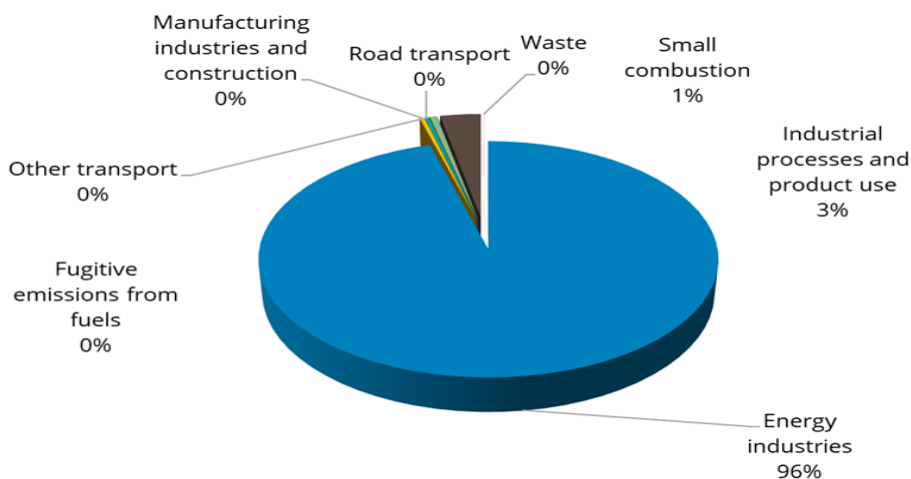


Figure 2.6.3.16 Individual sectors contribution of Se emissions for 2017

### Zinc Emissions

National zinc (Zn) emissions increased from the year 1990, when total amount was 18,5 t to 21,7 t in 2017. Emissions were increased between 1990 and 2017 by 17 %. Significant drop of emissions in 2014 was due to smaller use of fuels in energy and small combustion sector. The main source for Zn emissions in the year 2017 was small combustion sector with a share of 45 %, followed by road transport with 25 %.

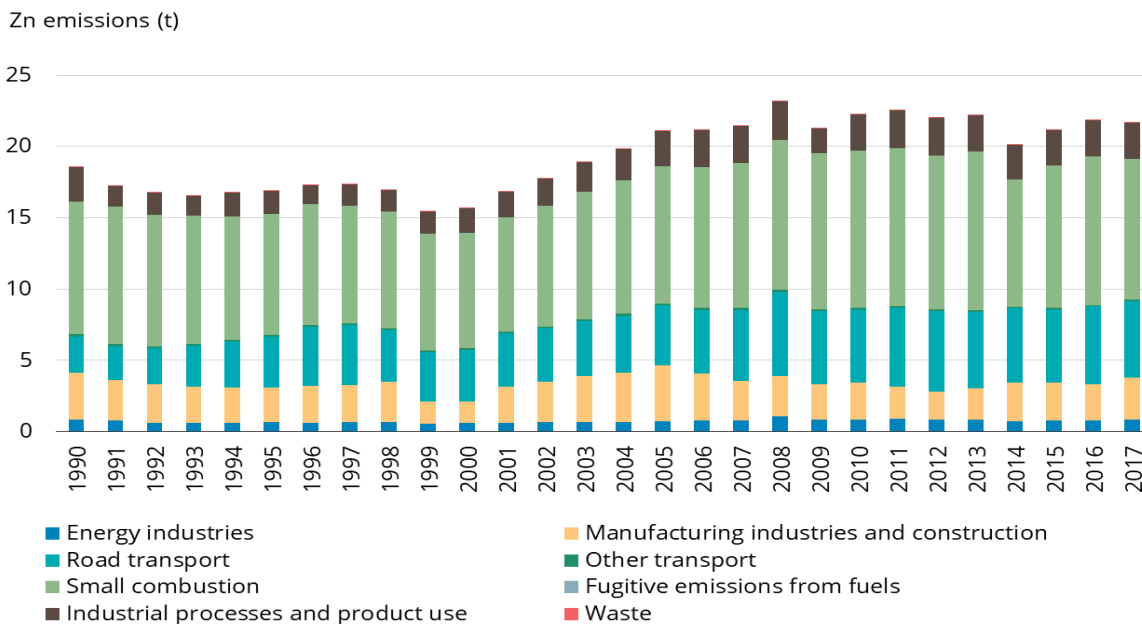


Figure 2.6.3.17 Zn emissions in Slovenia for the period 1990 –2017

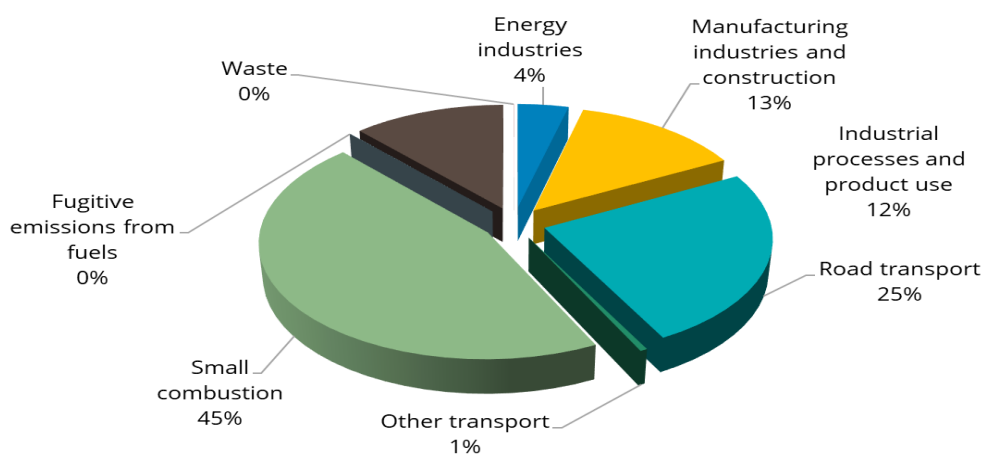


Figure 2.6.3.18 Individual sectors contribution of Zn emissions for 2017

#### 2.6.4 Emission Trends for Persistent Organic Pollutants

Persistent Organic Pollutants (POPs) is a common name of a group of pollutants that are semi-volatile, bioaccumulative, persistent and toxic. POPs are recognised as being directly toxic to biota. All have the quality of being progressively accumulated higher up the food chain, such that chronic exposure of lower organisms to much lower concentrations can expose predatory organisms, including humans and wildlife, to potentially harmful concentrations. In humans they are also of concern for human health because of their toxicity, their potential to cause cancer and their ability to cause harmful effects at low concentrations. Their relative toxic/carcinogenic potencies are compound specific. POPs including PAHs have also been shown to possess a number of toxicological properties. The major concern is centred on their possible role in carcinogenic, immunological and reproductive effects but more recently concern has also been expressed over their possible harmful effects on human development. The overall and long-term goal of the Aarhus Protocol on POPs is to eliminate any discharges, emissions and losses of POPs to the environment. Another agreement, which is ratified by Slovenia, is Stockholm Convention on Persistent Organic Pollutants. Within these conventions, the establishment of emission inventories for POPs is mandatory and provides the basis for further emission reductions among Parties.

In general, the most accurate way to establish emission rates is to measure them. However in most cases only limited measurements data are available. Therefore several guidebooks, guidelines and scientific literature make proposals for emission estimates when measurements data are lacking. In Slovenia emission national emission factors are not available; therefore they were taken from EMEP/EEA Emission inventory guidebook, 2016.

Persistent Organic Pollutants have been reported:

- Polycyclic aromatic hydrocarbons (PAHs): benzo(a)pyrene, benzo(k)fluoranthene, benzo(b)fluoranthene, indeno(1,2,3-cd)pyrene
- Dioxins and furans
- Hexachlorobenzene (HCB)
- Polychlorinated Biphenyls (PCB)

Emissions of PCB, dioxins and furans and PAH declined since 1990 as a result of decreased residential use of coal, improvements in abatement technologies for metal refining and smelting, and stricter regulations on emissions from the road transport sector. Implementation of legislation, stricter inspection and use of best available techniques has been responsible for decrease of POPs in last two decades. Emissions of HCB has increased in the same period due to bigger fuel consumption in transport sector and manufacturing industries and construction.

Emissions of POPs declined substantially from year 1990 to 2017: for PCB (91 %), dioxins/furans (15 %), PAH (35 %) and HCB (97 %)

Slovenia in 2017 did not exceed emission levels set in protocol on persistent organic pollutants for PCB, dioxins/furans, HCB and PAHs. Emissions are much below values from the reference year 1990.

Table 2.6.4.1 National total emissions and emission trends for PCB, dioxins/furans, PAHs and HCB for the period 1990 - 2017

Year	PCB	Dioxins/ furans	PAH	HCB
	kg		Total 1- 4	
	kg	g I-Teq	t	kg
1990	415,4	18,3	8,1	19,0
1991	414,2	18,0	8,8	16,6
1992	372,9	17,0	7,7	15,3
1993	349,3	16,1	6,9	15,4
1994	322,0	15,5	6,2	15,3
1995	290,3	15,3	5,9	15,3
1996	273,9	15,0	5,6	14,8
1997	255,1	14,8	5,3	15,4
1998	243,8	14,9	5,2	15,3
1999	227,2	14,7	5,2	15,7
2000	213,5	15,0	5,1	19,1
2001	201,8	15,4	5,1	21,2
2002	184,1	14,8	5,4	0,6
2003	154,2	15,5	5,6	0,6
2004	142,5	16,1	5,8	0,5
2005	134,7	16,6	6,0	0,6
2006	122,3	17,0	6,1	0,6
2007	99,3	17,3	6,2	0,6
2008	93,7	17,9	6,3	0,6
2009	82,5	17,0	6,1	0,6
2010	75,6	17,4	6,1	0,6
2011	50,7	17,6	6,1	0,6
2012	43,7	17,0	5,9	0,6
2013	40,5	17,1	6,0	0,6
2014	40,5	14,6	5,0	0,5
2015	38,9	15,4	5,3	0,5
2016	38,9	15,8	5,4	0,6
2017	35,6	15,5	5,2	0,5
<b>Reduction trend (%)</b>	<b>-91,4 %</b>	<b>-15,2 %</b>	<b>-35,1 %</b>	<b>-97,2 %</b>

The sum of emissions of four individual species: benzo(a)pyrene, benzo(k)fluoranthene, benzo(b)fluoranthene, indeno(1,2,3-cd)pyrene could be expressed as PAH Total 1-4 emission. In some cases emission factors for individual PAHs are not available, but there is an emission factor given only for Total 1-4. The sum of individual species does not always equal to Total 1-4 emission.

## PAH Emissions

Polycyclic aromatic hydrocarbons (PAHs) are a group of compounds composed of two or more fused aromatic rings and do not contain heteroatoms or carry substituents. The UNECE POPs Protocol specified that the following 4 PAHs should be used as indicators for the purposes of emission inventories: benzo(a)pyrene, benzo(k)fluoranthene, benzo(b)fluoranthene, indeno(1,2,3-cd)pyrene. PAH Total 1-4 emission is the sum of emissions of four individual species.

Table 2.6.4.2 PAHs emissions for the year 2017

Pollutant	Benzo(a) pyrene	Benzo(b) fluoranthene	Benzo(k) fluoranthene	Indeno (1,2,3-cd) pyrene	Total 1-4
Unit	t	t	t	t	t
Emissions	2,10	1,24	1,17	0,40	5,24

National PAH emissions decreased from 8,1 t in the year 1990 to 5,2 t in year 2017. Emissions were reduced by 35 %. The most significant emission source of PAH were residential combustion processes (open fires, coal and wood burning for heating purposes) with a share of 79 %. Emissions have declined since 1990 as a result of decreased residential use of coal and improvements in abatement technologies. The reason for increase of emissions in 2009 was bigger use of wood biomass in the residential sector. Increasing consumption of biomass is probably a result of economic crisis and a high price of petroleum products as well as state measures to promote renewable energy sources. The decrease in emissions in the last two years was due to significantly reduced emissions from residential combustion. Warmer winter and improved thermal insulation of buildings contributed to lower fuel consumption.

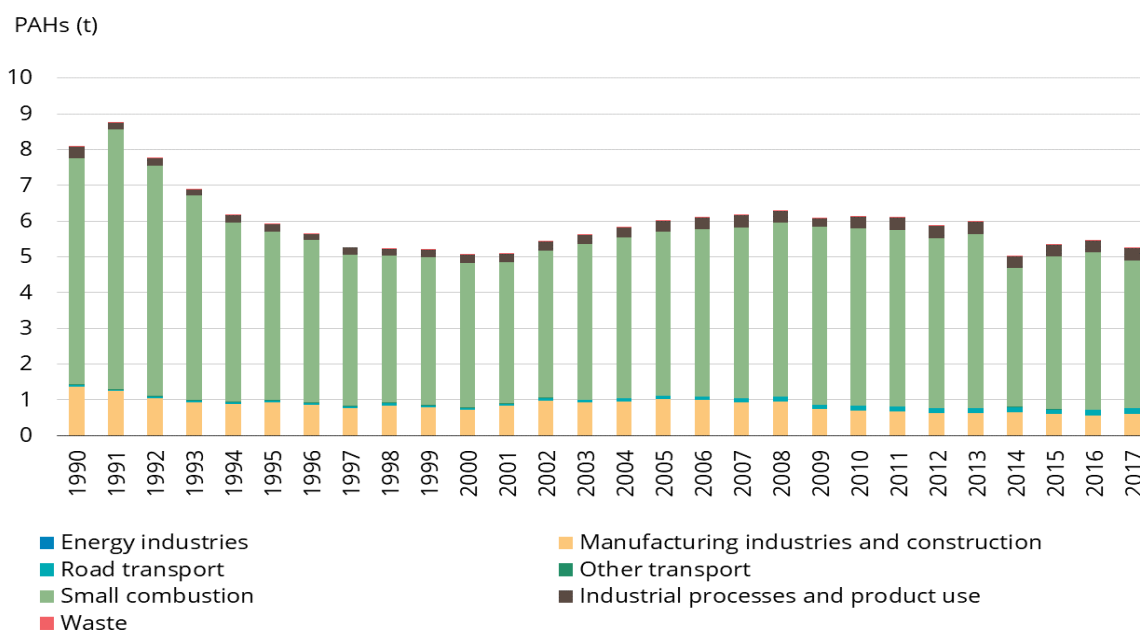


Figure 2.6.4.1 PAH emissions in Slovenia in the period 1990 – 2017

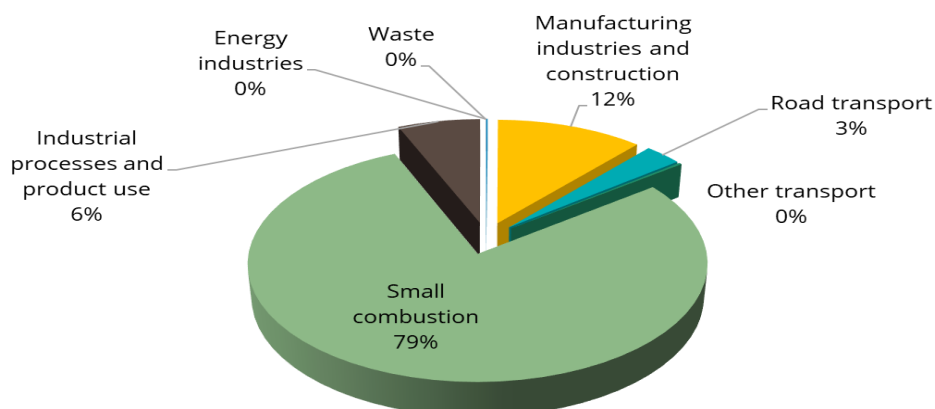


Figure 2.6.4.2 Individual sectors contribution of PAHs emissions for 2017

### PCB Emissions

National PCB emissions steadily decreased from the year 1990, when total amount was 415,4 kg to 35,6 kg in the year 2017. Emissions were reduced by 91 %, mainly due to reductions in product use subsector. Emissions have fallen due to phasing out of electrical equipment containing PCB. The main source for PCB emissions is industrial processes and product use with a share of more than 99 %.

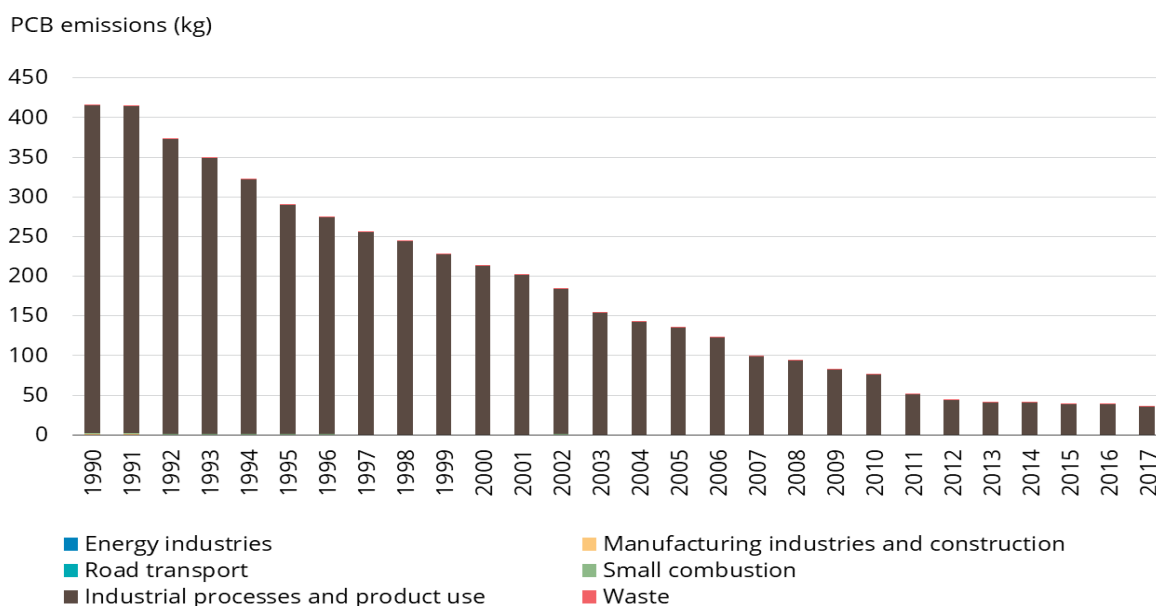


Figure 2.6.4.3 PCB emissions in Slovenia in the period 1990 –2017



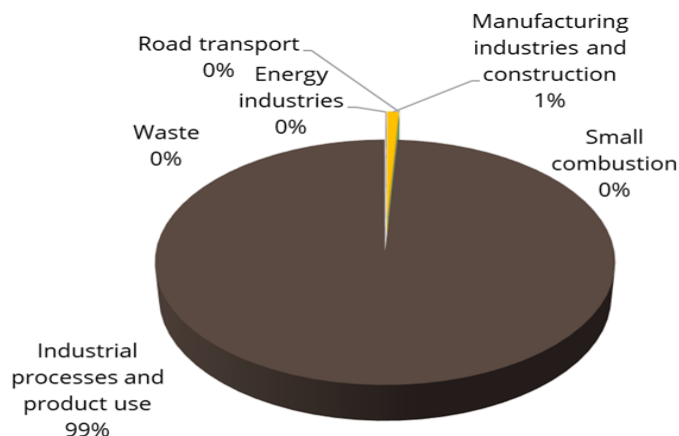


Figure 2.6.4.4 Individual sectors contribution of PCB emissions for 2017

### Dioxins and Furans Emissions

National dioxins and furans emissions steadily decreased from the year 1990, when total amount was 18,3 g I-TEQ to 15,5 g I-TEQ in 2017. Emissions were reduced by 15 %. The main sources of dioxins/furans emissions in 2017 were small combustion with a share of 62 % and industrial processes and product use with 15 %. The reason for increase of emissions in 2009 was bigger use of wood biomass in the residential sector. Increasing consumption of biomass is probably a result of economic crisis and a high price of petroleum products as well as state measures to promote renewable energy sources. The decrease in emissions in 2014 and 2015 was due to significantly reduced emissions from residential combustion. Warmer winter and improved thermal insulation of buildings contributed to lower fuel consumption.

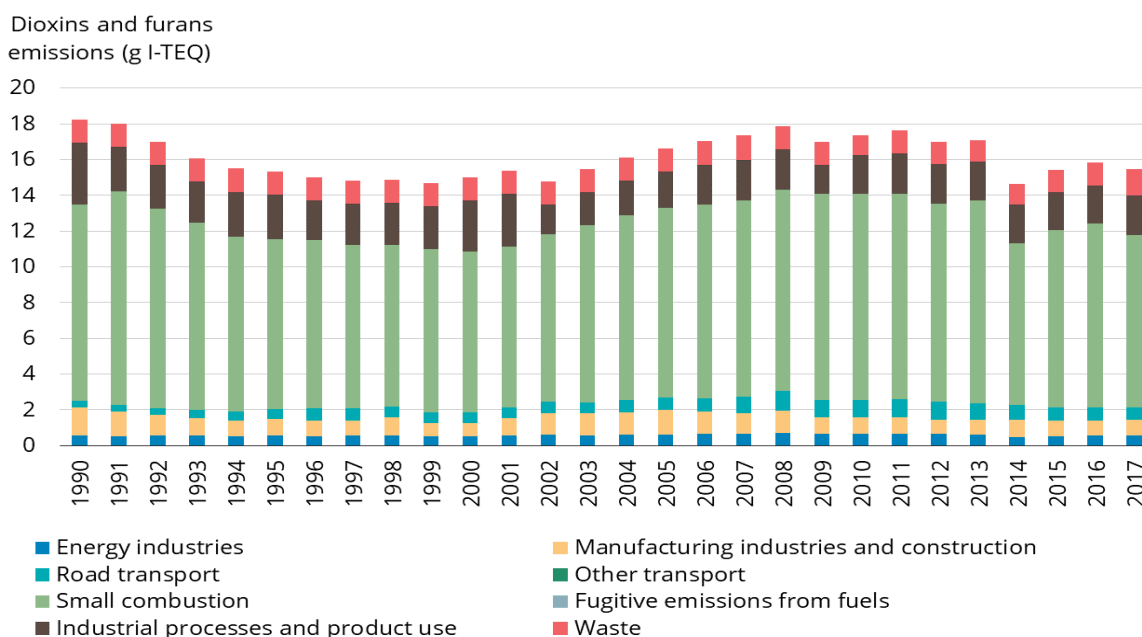


Figure 2.6.4.5 Dioxins and furans emissions in Slovenia for the period 1990 – 2017

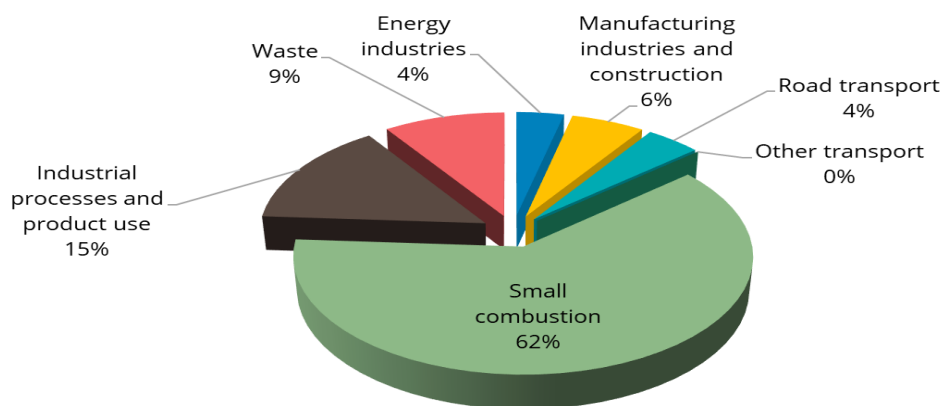


Figure 2.6.4.6 Individual sectors contribution of dioxins and furans emissions for 2017

### HCB Emissions

Emissions of HCB have decreased significantly since 1990 when total amount was 19,0 kg to 0,5 kg in 2017. Emissions were decreased by 97 %. The reason for drastic drop of emissions in 2002 was termination of hexachloroethane (HCE) tablets as a degassing agent in aluminium production.

In 2017, the main source for HCB emissions in Slovenia was heat and electricity production, with a share of 62 %, followed by small combustion sector (18 %).

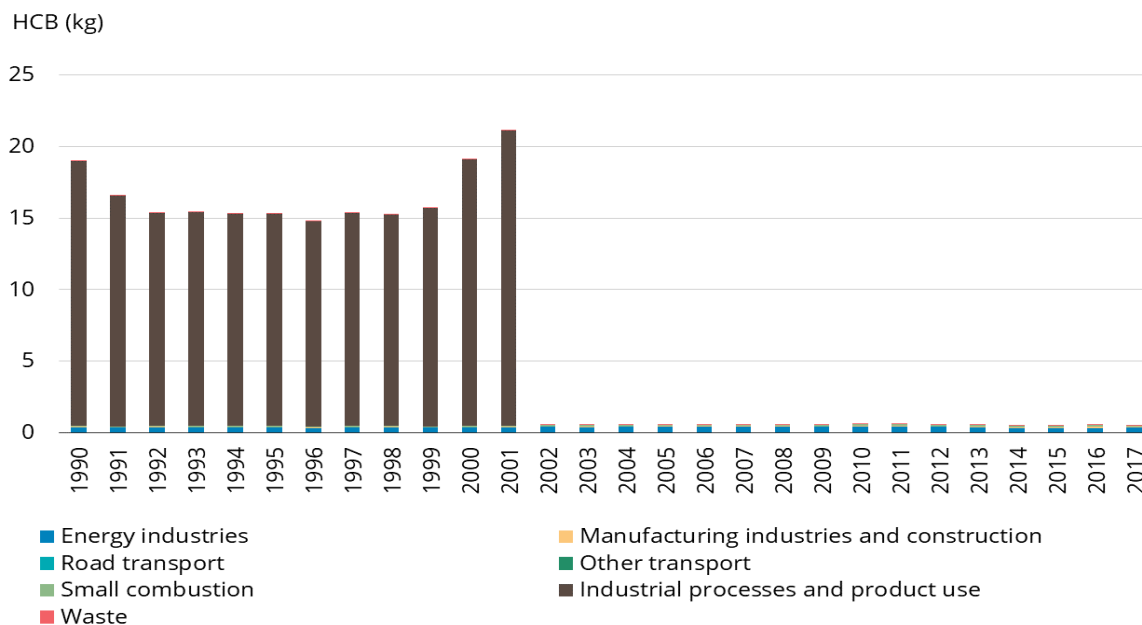


Figure 2.6.4.7 HCB emissions in Slovenia for the period 1990 – 2017

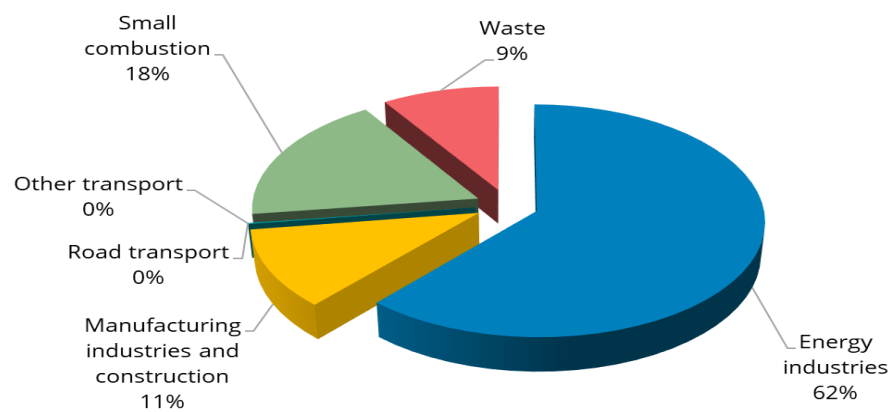


Figure 2.6.4.8 Individual sectors contribution of HCB emissions for 2017

### 3 ENERGY

The energy sector is the most important sector considering major air pollutants air emissions in the Republic of Slovenia. Emissions from this sector arise from fuel combustion (NFR sector 1. A) and fugitive emissions from fuels (NFR sector 1. B).

#### 3.1 Energy Industries (1. A. 1)

This chapter describes the methods and data needed to estimate emissions from NFR Sector 1A1 Energy industries. The activity covers combustion and conversion of fuels to produce energy, for example electricity or heat from point sources:

NFR Codes:

- 1A1a Public electricity and heat production
- 1A1b Petroleum refining
- 1A1c Manufacture of solid fuels

Public electricity and heat production is the most important category in this sub-sector. Other two categories consist mainly of fuel consumption in one refinery (closed in 2004) and in fuel consumption for coal mining activities and gas extraction.

##### 3.1.1 Public electricity and heat production

NFR Code 1A1a

Until 2015 there have been three big point sources in the Republic of Slovenia, which represented the backbone of the production of electrical energy from thermal power plants: Šoštanj Thermal Power Plant (TEŠ), Trbovlje Thermal Power Plant (TET) and Termoelektrarna Ljubljana (TE-TOL). All three plants have used coal for the production of electrical energy. Two of these thermal power plants, TEŠ and TET, are located beside coal pits. Since 2003, TE-TOL uses exclusively imported coal with high net calorific value and low sulphur contents for the production of electrical energy and heat.

In 2014, TET power plant was closed down. There are only two thermal power plants in operation since 2015.

**Table 3.1.1.1 Public electricity and Combined Heat and Power Plants in Slovenia**

Power plant	Location	Unit	Year	Power (MW)	Main fuel type
TEŠ	Šoštanj	A/1	1956-2010	30.0	Lignite from Velenje
TEŠ	Šoštanj	A/2	1956-2008	30.0	Lignite
TEŠ	Šoštanj	A/3	1960-2014	75.0	Lignite
TEŠ	Šoštanj	Unit 4	1972	275.0	Lignite
TEŠ	Šoštanj	Unit 5	1977	345.0	Lignite
TEŠ	Šoštanj	Unit 6	2016	600.0	Lignite
TEŠ	Šoštanj	Gas units	2008	2 x 42.0	Natural gas
TE-TOL	Ljubljana	D/1	1966	136.0	Imported coal
TE-TOL	Ljubljana	D/2	1967	126.0	Imported coal
TE-TOL	Ljubljana	D/3	1984	202.0	Imported coal, since 2008 also wood
TET	Trbovlje	F/4	1968-2014	125.0	Coal, mostly domestic brown coal

Besides thermal power plants we have also one small plant Brestanica – TEB which use natural gas and operate mainly as back up plant when more electricity is needed or when any other plant is on refit.

### Methodology

To estimate emissions from Public Electricity and Heat Production, the following methodologies have been adopted:

$$E = m \times \text{NCV} \times \text{EF} \quad \text{Equation 1}$$

E - emission (g)  
 m - quantity of fuel combusted (t)  
 NCV - net calorific value (TJ/kt)  
 EF - emission factor per energy of fuel (g/GJ)

$$E = m \times \text{EF} \quad \text{Equation 2}$$

E - emission (g)  
 m - quantity of fuel combusted (t)  
 EF - emission factor per quantity of fuel (g/t)

To estimate SO<sub>x</sub> emissions in same cases the following two equations for calculation of EF were used:

$$\text{EF}_{\text{SO}_x} = [\text{S}] \times 20000 / \text{NCV} \quad \text{Equation 3}$$

EF<sub>SO<sub>x</sub></sub> - SO<sub>x</sub> emission factor (g/GJ)  
 [S] – sulphur content of the fuel (% w/w)  
 NCV - net calorific value (GJ/t)  
 2 – ratio of the relative molecular mass of SO<sub>2</sub> to sulphur

$$\text{EF}_{\text{SO}_x} = [\text{S}] \times 19000 / \text{NCV} \quad \text{Equation 4}$$

EF<sub>SO<sub>x</sub></sub> - SO<sub>x</sub> emission factor (g/GJ)  
 [S] – sulphur content of the fuel (% w/w)  
 NCV - net calorific value (GJ/t)  
 1.9 – ratio of the relative molecular mass of SO<sub>2</sub> to sulphur, considering 5 % absorption in the ash

### Activity data

The main source of data for all energy industries in the Republic of Slovenia for the period 1980 - 2003 is LEG – Annual Energy Statistics of the Energy Sector of the Republic of Slovenia. As LEG was not published early enough to enable us to calculate national inventory on time in 2005 we have for the first time received data directly from Statistical Office of the Republic of Slovenia (SORS) in electronic format before they are published. These excel sheets are going to be our source of data for all fuel consumption in the future. Since 2005 all public power plants are included into ETS and verified reports from ETS have been used as data source.

Emissions from category “Other fuels” have arisen from Slovenian only waste incineration thermal plant which has started to work in 2009. Data on amount of incinerated waste, NCVs and distribution between biogenic and other waste have been obtained directly from the plant.

It shows up that the most of the waste in non biogenic part of waste is plastics. Because plastic is made from fossil fuels, its combustion is considered an anthropogenic source of carbon emissions.

Data on fuel consumption by type and year are reported in the Annex 1 to the IIR (Table 1.14: Fuel used in Energy industries 1980–2017).

### Net calorific values

Net calorific values (NCV) have been taken from SORS except for coal since 2005 when all three thermal power plants were included into the ETS and very detailed data on NCV become available. The values for solid fuel varies from year to year but for the liquid and gaseous fuel almost the same values have been used for the entire period as these types of fuel don't change a lot from year to year.

Table 3.1.1.2 NCVs for the fuel used in energy industry

Year	Lignite – domestic	Sub-bituminous Coal - domestic	Sub-bituminous Coal - imported	Residual Fuel Oil	Heavy Fuel Oil	Liquefied Petroleum Gas (LPG)	Natural Gas	Wood and Other Biomass	Waste
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/Mm3	TJ/kt	TJ/kt
1980	9,360	12,980		41,800	39,700		33,500	12,170	
1981	9,330	11,570		41,800	39,700		34,100	12,170	
1982	9,330	11,570		41,900	39,800		33,490	12,170	
1983	9,610	11,180		41,900	39,800		33,800	12,170	
1984	9,590	11,420		41,900	40,000		33,500	12,170	
1985	9,430	11,690		41,900	39,800		33,500	12,170	
1986	9,390	11,880		41,820	39,740	43,190	33,500	12,170	
1987	9,650	11,820		41,780	39,800	42,870	33,500	12,170	
1988	9,440	12,000		41,710	39,800	43,100	34,080	12,170	
1989	9,820	12,050		41,850	39,800	43,070	34,100	12,170	
1990	9,810	12,760		41,870	39,800	43,070	34,100	12,170	
1991	9,980	12,879		41,880	39,800	43,170	34,100	12,170	
1992	10,260	12,589		41,900	39,900	43,100	34,100	12,170	
1993	10,070	12,050		41,900	39,800	46,050	34,100	12,170	
1994	9,960	12,666		41,900	39,860	46,050	34,100	12,170	
1995	10,220	11,250	17,410	41,900	40,000	46,050	34,100	12,170	
1996	9,690	11,300	17,410	41,900	40,000	46,050	34,100	12,170	
1997	9,610	11,300	17,360	41,900	40,000	46,050	34,080	12,170	
1998	10,010	11,230	17,760	41,900	40,000	46,050	34,080	12,170	
1999	9,690	11,110	17,560	41,900	40,000	46,050	34,080	12,170	
2000	10,170	11,230	17,940	41,900	40,000	46,050	34,080	12,170	
2001	10,660	10,660	17,940	41,900	40,000	46,050	34,080	12,170	
2002	10,350	11,220	18,380	41,900	40,000	46,050	34,080	12,170	
2003	10,138	11,560	18,310	41,900	40,000	46,050	34,080	12,170	
2004	10,301	11,680	18,676	42,600	41,420	46,050	34,080	12,170	
2005	10,803	11,724	18,180	42,600	41,420	46,050	34,080	10,714	
2006	11,132	10,880	18,874	41,900	40,000	46,050	34,072	12,170	
2007	11,258	11,629	18,275	42,634	41,374	46,050	34,078	9,141	
2008	10,949	10,641	17,735	42,600	41,420	46,050	34,096	11,511	
2009	10,894	11,094	17,872	42,600	41,420	46,050	34,074	11,128	27,800

2010	11,097	12,815	18,130	42,600	41,420	46,050	34,080	9,871	27,800
2011	11,068	11,935	18,428	42,600	41,420	46,050	34,087	10,267	27,800
2012	10,616	11,778	18,524	42,600	41,420	46,050	34,093	10,559	27,800
2013	11,591	11,946	18,457	42,600	41,420	46,050	34,079	10,262	27,762
2014	10,823	11,727	18,655	42,600	41,420	46,050	34,083	10,510	27,762
2015	11,418	-	18,629	42,600	41,420	46,050	34,086	10,474	26,700
2016	11,733	-	18,595	42,600	41,420	46,050	34,087	10,519	26,700
2017	11,640	-	18,230	42,600	41,420	46,050	34,085	10,706	26,700

### Emission factors

County specific emissions factors were used for emission calculations of NO<sub>x</sub>, SO<sub>x</sub>, CO and particulate matter for the period 1980 – 2008 for domestic lignite, domestic sub-bituminous coal and imported sub-bituminous coal. Country specific emission factors were obtained from Electro Institute Milan Vidmar.

For the period 2009–2017 direct emissions have been taken from REMIS database, established and handled by Slovenian Environmental Agency. These data represent plant specific values.

REMIS database is obtained in compliance with Rules on initial measurements and operational monitoring of the emission of substances into the atmosphere from the stationary pollution sources and on the conditions for their implementation (OJ RS, No. 105/08). Each year all obligators must provide report on implementation of emission monitoring of substances into air. Annual emission report includes emissions of substances into air. These emissions data are direct measurements of emissions into air and reflect plant specific values.

According to 2017 in-depth EU NECD review thorough examination of annual emissions reported by operators was performed. All operators were checked individually. We carried out a survey for each company and we eliminated the risk of misinterpretation of measurement data. It was confirmed that the values that we used for the estimation of national emissions are not validated average values with the confidence limits subtracted. Reported data in Slovenian national inventory are raw measured values. Data used for NECD and CLRTAP reporting are not processed or changed in any way. The national emissions are not underestimated.

The validated average values where confidence interval is subtracted are used for other purpose, this is for determination of exceeding the emission limit values. Those data are not used for reporting of national emissions.

**Table 3.1.1.3 National emission factors for NO<sub>x</sub>, SO<sub>x</sub>, CO, PM<sub>2.5</sub>, PM<sub>10</sub>, TSP for domestic lignite from Velenje pit until 2008**

Year/ polutant	NO <sub>x</sub>	SO <sub>x</sub>	CO	PM <sub>10</sub>	PM <sub>2.5</sub>	TSP
Unit	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ
1980	364,85	2638,89	13,78			
1981	368,97	2647,37	14,45			
1982	356,81	2647,37	13,31			
1983	346,68	2570,24	12,84			
1984	349,12	2575,60	13,01			
1985	342,26	2619,30	12,83			
1986	344,39	2630,46	12,57			
1987	363,89	2559,59	13,48			
1988	351,48	2616,53	12,82			
1989	372,76	2515,27	14,20			

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1990	346,05	2517,84	13,19			
1991	319,35	2474,95	12,93			
1992	271,16	2407,41	13,04			
1993	292,99	2452,83	13,22			
1994	314,32	2479,92	13,41			
1995	269,89	1378,66	20,29			
1996	295,55	1489,82	18,19			
1997	298,06	1367,70	19,01			
1998	290,92	1339,51	17,86			
1999	251,85	1319,67	16,26			
2000	273,86	1170,24	14,26	9,123	4,257	12,164
2001	268,50	425,71	16,31	8,251	3,851	11,002
2002	283,91	508,67	20,69	10,542	4,920	14,056
2003	264,14	322,49	24,98	8,707	4,063	11,609
2004	206,29	184,91	30,21	7,308	3,411	9,744
2005	208,61	238,46	19,79	5,742	2,680	7,656
2006	205,27	139,30	18,59	2,667	1,244	3,556
2007	183,93	115,12	27,33	3,415	1,594	5,533
2008	188,61	103,87	23,20	3,664	1,710	4,886

**Table 3.1.1.4 National emission factors for NO<sub>x</sub>, SO<sub>x</sub>, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP for domestic sub-bituminous from Trbovlje coalmine until 2008**

Year/ polutant	NO <sub>x</sub>	SO <sub>x</sub>	PM <sub>10</sub>	PM <sub>2.5</sub>	TSP
Unit	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ
1980	225,86	2927,58			
1981	226,21	3284,36			
1982	233,71	3284,36			
1983	238,61	3398,93			
1984	242,16	3327,50			
1985	265,12	3250,64			
1986	231,83	3198,65			
1987	235,22	3214,89			
1988	231,65	3166,67			
1989	199,05	3153,53			
1990	212,25	2978,06			
1991	185,24	2950,45			
1992	220,48	3018,57			
1993	237,27	3153,53			
1994	223,03	3000,16			
1995	192,96	3377,78			
1996	201,32	3867,26			
1997	216,58	4203,54			
1998	190,01	4229,74			
1999	253,21	4275,43			
2000	247,92	4229,74	36,529	17,047	48,706
2001	187,97	4099,44	35,908	16,757	47,878
2002	239,31	3894,83	34,700	26,000	39,232
2003	233,06	4602,08	34,281	15,998	45,708
2004	282,08	4554,79	41,526	19,379	55,368
2005	243,15	3076,35	39,796	18,571	53,061



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2006	235,43	284,07	7,507	3,503	10,009
2007	197,54	296,93	10,145	4,734	13,527
2008	190,00	289,40	15,991	7,463	21,322

**Table 3.1.1.5 National emission factors for NO<sub>x</sub>, PM<sub>2.5</sub>, PM<sub>10</sub>, TSP and sulphur content for imported sub-bituminous coal until 2008**

Year/ pollutant	NO <sub>x</sub>	PM <sub>10</sub>	PM <sub>2.5</sub>	TSP	SO <sub>x</sub> Equation 4
Unit	g/GJ	g/GJ	g/GJ	g/GJ	[S] (% w/w)
1990					
1991					
1992					
1993					
1994					
1995	200,00				1,60
1996	220,00				1,60
1997	280,00				1,60
1998	280,00				0,12
1999	230,00				0,12
2000	210,00	8,000	6,000	9,000	0,12
2001	220,00	8,000	6,000	9,000	0,12
2002	190,00	13,648	6,369	18,197	0,07
2003	180,00	6,460	3,015	8,613	0,09
2004	164,02	6,246	2,915	8,328	0,09
2005	162,97	6,994	3,264	9,326	0,14
2006	177,38	6,090	2,842	8,119	0,14
2007	154,61	2,539	1,185	3,386	0,14
2008	156,86	3,554	1,659	4,739	0,10

In calculating emissions of other individual gases, following emission factors have been used:

**Table 3.1.1.6 Emission factors used for domestic lignite and domestic sub-bituminous coal for the period 1990 - 2017**

Pollutant	Value	Unit	References
NM VOC	1,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Cd	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Pb	15	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Hg	2,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
As	14,3	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Cr	9,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Cu	1,0	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Ni	9,7	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Se	45	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17

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Zn	8,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Dioxins/ Furans	10	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Benzo(a)pyrene	1,3	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Benzo(b)fluoranthene	37	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Benzo(k)fluoranthene	29	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Indeno(1,2,3-cd)pyrene	2,1	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
HCB	6,7	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
CO	8,7 (except for domestic lignite, see Table 3.1.1.3)	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17

Table 3.1.1.7 Emission factors used for imported sub-bituminous coal for the period 1995 - 2017

Pollutant	Value	Unit	References
NM VOC	1,0	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Cd	0,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Pb	7,3	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Hg	1,4	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
As	7,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Cr	4,5	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Cu	7,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Ni	4,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Se	23	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Zn	19	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Dioxins/ Furans	10	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(a)pyrene	0,7	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(b)fluoranthene	37	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(k)fluoranthene	29	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Indeno(1,2,3-cd)pyrene	1,1	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
HCB	6,7	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16

Emission factor for Hg was corrected for domestic lignite and domestic sub-bituminous coal. Correction of EF was performed due to use of flue-gas desulfurization device. Prescribed emission factor without flue-gas desulfurization applied is 2,9 mg/GJ. Estimation of Hg capture by currently installed pollution control equipment range from 47-81 % Hg capture for electrostatic precipitators and flue-gas desulfurization.

Table 3.1.1.8 Emission factors used for heavy fuel oil for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	142	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
SO <sub>x</sub>	Equation 3	[S] (% w/w), see Table 3.1.1.9	Slovene national legislation relating quality of liquid fuels
CO	15,1	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
NM VOC	2,3	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
PM <sub>10</sub>	25,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
PM <sub>2.5</sub>	19,3	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
TSP	35,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
BC	1,081	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cd	1,2	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Pb	4,56	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Hg	0,341	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
As	3,98	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cr	2,55	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cu	5,31	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Ni	255	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Se	2,06	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Zn	87,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Dioxins/ Furans	2,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Benzo(b)fluoranthene	4,5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Benzo(k)fluoranthene	4,5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Indeno(1,2,3-cd)pyrene	6,92	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19

Table 3.1.1.9 Emission factors used for residual fuel oil for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	65	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
SO <sub>x</sub>	Equation 3	[S] (% w/w), see Table 3.1.1.9	Slovene national legislation relating quality of liquid fuels
CO	16,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
NM VOC	0,8	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20

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<b>PM<sub>10</sub></b>	3,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>PM<sub>2.5</sub></b>	0,8	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>TSP</b>	6,5	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>BC</b>	0,268	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cd</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Pb</b>	4,07	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Hg</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>As</b>	1,81	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cr</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cu</b>	2,72	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Ni</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Se</b>	6,79	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Zn</b>	1,81	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Dioxins/ Furans</b>	0,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Indeno(1,2,3-cd)pyrene</b>	6,92	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20

Table 3.1.1.10 Sulphur content in residual fuel oil and heavy fuel oil for 1980 - 2017

<b>Fuel</b>	<b>Heavy fuel Oil</b>	<b>Residual fuel Oil</b>	<b>Fuel</b>	<b>Heavy fuel Oil</b>	<b>Residual fuel Oil</b>
<b>Year</b>	<b>[S] (% w/w)</b>	<b>[S] (% w/w)</b>	<b>year</b>	<b>[S] (% w/w)</b>	<b>[S] (% w/w)</b>
1980	3,0	1,2	1999	1,0	0,2
1981	3,0	1,2	2000	1,0	0,2
1982	3,0	1,2	2001	1,0	0,2
1983	3,0	1,2	2002	1,0	0,2
1984	3,0	1,2	2003	1,0	0,2
1985	3,0	1,2	2004	1,0	0,2
1986	3,0	1,2	2005	1,0	0,2
1987	3,0	1,2	2006	1,0	0,2
1988	3,0	1,2	2007	1,0	0,2
1989	3,0	1,2	2008	1,0	0,1
1990	3,0	1,2	2009	1,0	0,1
1991	3,0	1,2	2010	1,0	0,1
1992	3,0	1,2	2011	1,0	0,1
1993	3,0	1,2	2012	1,0	0,1
1994	3,0	1,2	2013	1,0	0,1
1995	1,5	0,5	2014	1,0	0,1
1996	1,0	0,2	2015	1,0	0,1
1997	1,0	0,2	2016	1,0	0,1
1998	1,0	0,2	2017	1,0	0,1

Table 3.1.1.11 Emission factors used for natural gas, biogas and liquefied petroleum gas for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
CO	39	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
SO <sub>x</sub>	0,281	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
NM VOC	2,6	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
PM <sub>10</sub>	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
PM <sub>2.5</sub>	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
TSP	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
BC	0,0223	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cd	0,00025	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Pb	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Hg	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
As	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cr	0,00076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cu	0,000076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Ni	0,00051	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Se	0,0112	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Zn	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Dioxins/ Furans	0,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(a)pyrene	0,56	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(b)fluoranthene	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(k)fluoranthene	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Indeno(1,2,3-cd)pyrene	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18

Table 3.1.1.12 Emission factors used for wood and other biomass for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	81	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
CO	90	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
NM VOC	7,31	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
SO <sub>x</sub>	10,8	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
PM <sub>10</sub>	155	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20

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<b>PM<sub>2.5</sub></b>	133	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>TSP</b>	172	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>BC</b>	4,389	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Cd</b>	1,76	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Pb</b>	20,6	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Hg</b>	1,51	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>As</b>	9,46	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Cr</b>	9,03	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Cu</b>	21,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Ni</b>	14,2	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Se</b>	1,2	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Zn</b>	181	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Benzo(a)pyrene</b>	1,12	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Benzo(b)fluoranthene</b>	0,043	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Benzo(k)fluoranthene</b>	0,0155	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Indeno(1,2,3-cd)pyrene</b>	0,0374	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>Dioxins/ Furans</b>	50	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>PCB</b>	3,5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20
<b>HCB</b>	5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-7, pg 20

Table 3.1.1.13 Emission factors used for waste 2009 - 2017

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	0,87	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>SO<sub>x</sub></b>	0,047	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>CO</b>	0,07	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>NM VOC</b>	7,4	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>PM<sub>2.5</sub></b>	0,004	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>PM<sub>10</sub></b>	0,007	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>TSP</b>	0,01	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>BC</b>	0,00014	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>Cd</b>	0,1	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>Hg</b>	0,056	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10

<b>Pb</b>	1,3	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>As</b>	0,016	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>Ni</b>	0,14	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>Dioxins/ Furans</b>	1	microg I-TEQ/t	Plant specific
<b>Total 4 PAHs</b>	0,02	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10
<b>HCB</b>	0,002	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, Table 3-1, pg 10

Data on particulate matter from hard and brown coal, heavy fuel oil and gas oil, biomass and gaseous fuel include filterable emissions. But there is no information whether PM emission factors used for waste include or exclude condensable component.

### Emissions

Public electricity and heat production is important source of SO<sub>x</sub> emissions. It contributed more than 40 % to total national emissions in 2017. It was even bigger SO<sub>x</sub> polluter before introduction of flue gas desulphurization device and gas turbines in power cogeneration plants. Emissions of most pollutants have decreased in last decades due to improvement in technologies, implementation of abatement techniques and fuel switching to cleaner fuels.

### Recalculations

According to 2018 in-depth EU NECD review imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 were used for emissions calculation. Recalculation of emissions were therefore performed. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb, NMVOC were recalculated for the period 1990-2016. Recalculation of PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC, SO<sub>x</sub> and NO<sub>x</sub> emissions were performed for the period 2009-2016.

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-specific QA/QC and verification

In 2005, all thermal power plants in the Republic of Slovenia have carried out regular coal sampling and determined the carbon contents in accordance with the Monitoring guidelines for monitoring and reporting of greenhouse gas emissions pursuant to Directive 2003/87/EC of European Parliament and of the Council and all amending directive, necessary for CO<sub>2</sub> emission trading on the territory of the European Union. The monitoring of fuel in four plants under EU-ETS is defined in the permit and accompanied monitoring plan. Each fuel is monitored with maximum uncertainty which depends on total GHG emissions from the plant and typical consumption of a particular fuel. All three plants have to monitor the coal consumption on the higher level of accuracy and determine NCV and carbon content in the accredited laboratory for every batch of fuel. The fourth plant is using natural gas as a main fuel.

For three thermal power plants the aggregated solid fuel from SORS data are compared with the sum of fuel used from verified ETS reports. The NCV values are also checked. In case these numbers are not the same as in ETS, data from ETS is taken into account and notification to SORS is made.

Additional QA activity is reference approach. Before entering data into database, the sum of

each fuel from disaggregated data is compared with energy balance data, reported in the Joint Questioner. As data in JQ are rounded to 1000 units, the difference should be 500 units or less. If it is higher, the reasons for this should be found.

According to 2017 in-depth EU NECD review thorough examination of annual emissions reported by operators was performed. All operators were checked individually. We carried out a survey for each company and we eliminated the risk of misinterpretation of measurement data. It was confirmed that the values that we used for the estimation of national emissions are not validated average values with the confidence limits subtracted. Reported data in Slovenian national inventory are raw measured values. Data used for NECD and CLRTAP reporting are not processed or changed in any way. The national emissions are not underestimated.

Information on condensable component of particulate matter was introduced into IIR 2019. A table summarising whether PM<sub>10</sub> and PM<sub>2.5</sub> emission factors for each source sector include or exclude the condensable component and references for their emission factors are presented in Annex 2 to the IIR 2019.

In addition, notation keys were revised as well. NFR tables were checked and corrected, if necessary.

### **Future improvements**

No improvement is planned for next submission.

### **3.1.2 Petroleum Refining**

NFR Code 1A1b

The main representative of this category was company the Nafta Lendava Refinery – Slovenian only refinery which stopped oil refining in 2002. According to the statistical methodology in the period 1986-1996, this sector also included quantities of fuels that were consumed for the production of electric energy in this sector.

Emissions of all pollutants from this sector were insignificant in the period 1980-2003. Since the only petroleum refinery was closed in 2003, no emissions have occurred from this category after 2003. Notation key “NO” (not occurring) have been used since 2004 for this sector.

### **Methodology**

To estimate emissions from Petroleum Refining, the same methodology as in Energy Industries was used.

### **Activity data**

Data on the consumption of fuels in this sector for the period 1986-2003 have been collected in LEG – Annual Energy Statistics of the Energy Sector of the Republic of Slovenia: for the period 1986-1996 under „Oil Industry”. From 1997 – 2004 under „DF–Production of coke, refined



petroleum products and nuclear fuel”.

- For the consumption of liquid fuels           Table Tg/3 or Table Pg/6 for LPG
- For the consumption of solid fuels           Table Pr/6
- For the consumption of gaseous fuels       Table Pg/6

After 1996, data on the consumption in this sector have been included in the industrial sector DF – Production of coke, refined petroleum products, and nuclear fuel. With regard to the fact there is neither production of coke nor nuclear fuel in the Republic of Slovenia, data for the period 1997-2003 are comparable to the data from the period 1986-1996. Data for the period 1980-1985 have been estimated.

Data on fuel consumption by type and year are reported in the Annex 1 to the IIR (Table 1.14: Fuel used in Energy industries 1980–2017).

### Net calorific values

Net calorific values have been taken from Statistical Office of the Republic of Slovenia.

**Table 3.1.2.1 NCVs for the fuel used in petroleum refining**

Year	Residual Fuel Oil	Heavy Fuel Oil	Natural gas
	TJ/kt	TJ/kt	TJ/Mm3
1980	41,82	39,74	33,50
1981	41,82	39,74	33,50
1982	41,82	39,74	33,50
1983	41,82	39,74	33,50
1984	41,82	39,74	33,50
1985	41,82	39,74	33,50
1986	41,82	39,74	33,50
1987	41,78	39,80	33,50
1988	41,71	39,80	34,08
1989	41,85	39,80	34,10
1990	41,87	39,80	34,10
1991	41,88	39,80	34,10
1992	41,90	39,90	34,10
1993	41,90	39,80	34,10
1994	41,90	39,86	34,10
1995	41,90	40,00	34,10
1996	41,90	40,00	34,10
1997	41,90	40,00	34,08
1998	41,90	40,00	34,08
1999	41,90	40,00	34,08
2000	41,90	40,00	34,08
2001	41,90	40,00	34,08
2002	41,90	40,00	34,08
2003	41,90	40,00	34,08

### Emission factors

For calculating emissions of individual gases in petroleum refining following emission factors have been used:

Table 3.1.2.2 Emission factors used for heavy fuel oil for 1980 – 2003

Pollutant	Value	Unit	References
NO <sub>x</sub>	142	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
SO <sub>x</sub>	Equation 3	[S] (% w/w), see Table 3.1.1.9	Slovene national legislation relating quality of liquid fuels
CO	15,1	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
NM VOC	2,3	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
PM <sub>10</sub>	25,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
PM <sub>2.5</sub>	19,3	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
TSP	35,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
BC	1,081	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cd	1,2	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Pb	4,56	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Hg	0,341	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
As	3,98	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cr	2,55	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Cu	5,31	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Ni	255	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Se	2,06	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Zn	87,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Dioxins/ Furans	2,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Benzo(b)fluoranthene	4,5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Benzo(k)fluoranthene	4,5	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19
Indeno(1,2,3-cd)pyrene	6,92	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-5, pg 19

Table 3.1.2.3 Emission factors used for residual fuel oil for 1980 - 2003

Pollutant	Value	Unit	References
NO <sub>x</sub>	65	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
SO <sub>x</sub>	Equation 3	[S] (% w/w), see Table 3.1.1.9	Slovene national legislation relating quality of liquid fuels
CO	16,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20

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<b>NMVOG</b>	0,8	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>PM<sub>10</sub></b>	3,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>PM<sub>2.5</sub></b>	0,8	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>TSP</b>	6,5	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>BC</b>	0,268	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cd</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Pb</b>	4,07	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Hg</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>As</b>	1,81	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cr</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Cu</b>	2,72	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Ni</b>	1,36	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Se</b>	6,79	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Zn</b>	1,81	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Dioxins/ Furans</b>	0,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20
<b>Indeno(1,2,3-cd)pyrene</b>	6,92	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-6, pg 20

Table 3.1.2.4 Emission factors used for natural gas for 1980 - 2001

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>CO</b>	39	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>SO<sub>x</sub></b>	0,281	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>NMVOG</b>	2,6	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>PM<sub>10</sub></b>	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>PM<sub>2.5</sub></b>	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>TSP</b>	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>BC</b>	0,0223	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Cd</b>	0,00025	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Pb</b>	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Hg</b>	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>As</b>	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18

<b>Cr</b>	0,00076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Cu</b>	0,000076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Ni</b>	0,00051	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Se</b>	0,0112	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Zn</b>	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Benzo(a)pyrene</b>	0,56	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Benzo(b)fluoranthene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Benzo(k)fluoranthene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Indeno(1,2,3-cd)pyrene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Dioxins/ Furans</b>	0,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors represent filterable PM emissions.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2003.

### Category-specific QA/QC and verification

The source category QA/QC for this sector was performed as explained in Public electricity and heat production sector.

### Future improvements

No improvements are planned for next submission.

### 3.1.3 Manufacture of solid fuels and Other energy Industries

NFR Code 1A1c

This sector covers the consumption of fuels reported in LEG under “Coal-mining” or, since 1997, under CA – Production of energy commodities and DF – Production of fuels.

Emissions of all pollutants from this sector are insignificant. This sector contributed in 2017 less than 0,05 % to total national emissions.

## Methodology

To estimate emissions from Manufacture of solid fuels and Other energy Industries the same methodology as in Energy Industries was used.

## Activity data

Consumptions according to individual energy products are collected in LEG tables as follows: For the period 1986-1996 under „Coal-mining”.

From 1997 onwards under „CA–Production of energy commodities”.

- For the consumption of liquid fuels Table Tg/3 or Table Pg/6 for LPG
- For the consumption of solid fuels Table Pr/6
- For the consumption gaseous fuels Table Pg/6

Since 2004, data are available in the excel files from SORS (E\_PE-M YYYY.xls).

In the period 2004 -2007 according to the old SKD classification the following SKD categories have been included in this CRF category:

- CA10 Mining of coal and lignite
- CA11 Extraction of crude petroleum and natural gas including support activities
- DF Production of coke, refined petroleum products and nuclear fuel

Since 2008, the new SKD\_2008 classification has been used and the following categories have been included in this CRF category:

- B05 Mining of coal and lignite
- B06 Extraction of crude petroleum and natural gas
- B09.1 Support activities for petroleum and natural gas mining
- C19.1 Manufacturing of coke oven products - do not exist in Slovenia.
- C19.2 Manufacturing of refined petroleum products

In the year 2017 only natural gas was consumed in this sector. Data on fuel consumption by type and year are reported in the Annex 1 to the IIR (Table 1.14: Fuel used in Energy industries 1980–2017).

## Net calorific values

Net calorific values have been taken from Statistical Office of the Republic of Slovenia.

**Table 3.1.3.1 NCVs and % S for the fuel used in Manufacture of solid fuels and other energy Industries**

Year	Sub-bituminous Coal - domestic	Sub-bituminous Coal - imported	Residual Fuel Oil	Heavy Fuel Oil	LPG	Natural Gas	Sub-bituminous Coal - domestic	Sub-bituminous Coal - imported
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/Mm3	[S] (% w/w)	[S] (% w/w)
1986	11,88		41,82		46,00	33,500	1,600	
1987	11,82		41,78		46,00	33,500	1,600	
1988	12,00		41,71		46,00	34,080	1,600	
1989	12,05		41,85		46,00	34,100	1,600	
1990	12,76		41,87		46,00	34,100	1,600	
1991	12,88		41,88		46,00		1,600	
1992	12,59		41,90	39,90	46,00	34,100	1,600	
1993	13,35		41,90		46,00	34,100	1,600	

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1994	12,67		41,90		46,00		1,600	
1995		17,40	41,90		46,00	34,100		1,600
1996		16,35	41,90		46,00			1,600
1997		17,71			46,05			1,600
1998		20,66	41,90					0,120
1999		20,81	41,90					0,120
2000		20,78	41,90					0,120
2001		20,95	41,90					0,120
2002			41,90					
2003			41,90					
2004			41,90		46,05			
2005			42,60	41,42	46,05			
2006			41,90	40,00	46,05	34,080		
2007			42,61	41,42	46,11			
2008			42,60	41,12	46,05	34,096		
2009			42,60			34,080		
2010			42,60			34,080		
2011			42,60			34,087		
2012			42,60			34,093		
2013			42,60			34,079		
2014						34,083		
2015						34,086		
2016						34,087		
2017						34,085		

### Emission factors

For calculating emissions of individual gases in manufacture of solid fuels and other energy industries emission factors used for residual fuel oil, heavy fuel oil and natural gas are the same as stated in chapter petroleum refining (Tables 3.1.2.2 - 3.1.2.4). Emission factors used for domestic sub-bituminous coal, imported sub-bituminous coal and liquefied petroleum gas are presented in the Tables 3.1.3.2, 3.1.3.3. and 3.1.3.4.

Table 3.1.3.2 Emission factors used for domestic sub-bituminous coal for 1986 - 1994

Pollutant	Value	Unit	References
NO <sub>x</sub>	247	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
SO <sub>x</sub>	Equation 4	[S] (% w/w, See Table 3.1.3.1	Slovene national legislation relating quality of liquid fuels
CO	8,7	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
NM VOC	1,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
PM <sub>10</sub>	7,9	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
PM <sub>2.5</sub>	3,2	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
TSP	11,7	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
BC	0,032	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
Cd	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17

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<b>Pb</b>	15	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Hg</b>	2,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>As</b>	14,3	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Cr</b>	9,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Cu</b>	1,0	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Ni</b>	9,7	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Se</b>	45	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Zn</b>	8,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Dioxins/ Furans</b>	10	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Benzo(a)pyrene</b>	1,3	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Benzo(b)fluoranthene</b>	37	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Benzo(k)fluoranthene</b>	29	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Indeno(1,2,3-cd)pyrene</b>	2,1	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>HCB</b>	6,7	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17

Table 3.1.3.3 Emission factors used for imported sub-bituminous coal for 1995 – 2001

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	209	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>SO<sub>x</sub></b>	<i>Equation 4</i>	[S] (% w/w, See Table 3.1.3.1	Slovene national legislation relating quality of liquid fuels
<b>CO</b>	8,7	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>NMVOC</b>	1,0	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>PM<sub>10</sub></b>	7,7	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>PM<sub>2.5</sub></b>	3,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>TSP</b>	11,4	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>BC</b>	0,0748	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-3, pg 17
<b>Cd</b>	0,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>Pb</b>	7,3	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>Hg</b>	1,4	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>As</b>	7,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>Cr</b>	4,5	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
<b>Cu</b>	7,8	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16

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Ni	4,9	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Se	23	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Zn	19	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Dioxins/ Furans	10	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(a)pyrene	0,7	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(b)fluoranthene	37	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Benzo(k)fluoranthene	29	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
Indeno(1,2,3-cd)pyrene	1,1	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16
HCB	6,7	microg /GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-2, pg 16

Table 3.1.3.4 Emission factors used for liquefied petroleum gas for 1986 - 2008

Pollutant	Value	Unit	References
NOx	89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
CO	39	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
SOx	0,281	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
NM VOC	2,6	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
PM10	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
PM2.5	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
TSP	0,89	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
BC	0,0223	g/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cd	0,00025	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Pb	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Hg	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
As	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cr	0,00076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Cu	0,000076	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Ni	0,00051	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Se	0,0112	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Zn	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(a)pyrene	0,56	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(b)fluoranthene	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
Benzo(k)fluoranthene	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18



<b>Indeno(1,2,3-cd)pyrene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18
<b>Dioxins/ Furans</b>	0,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Energy industries, Table 3-4, pg 18

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors represent filterable PM emissions.

### Recalculations

According to 2018 in-depth EU NECD review imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 were used for emissions calculation. Recalculation of emissions were therefore performed. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb, NMVOC and NO<sub>x</sub> were recalculated for the period 1995-2001. Recalculation of PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC emissions were performed for 2000 and 2001.

Due to new activity data on natural gas obtained for the year 2016 recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene and indeno(1,2,3-cd)pyrene emissions was performed for the year 2016.

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-specific QA/QC and verification

The source category QA/QC for this sector was performed as explained in Public electricity and heat production sector.

### Future improvements

No improvements are planned for next submission.

## 3.2 Manufacturing Industries and Construction (1. A. 2)

### 3.2.1 Stationary Combustion in manufacturing industries and construction

Sectors covered in this chapter are:

NFR Codes:

1A2a	Stationary combustion in manufacturing industries and construction: Iron and steel
1A2b	Stationary combustion in manufacturing industries and construction: Non-ferrous metals
1A2c	Stationary combustion in manufacturing industries and construction: Chemicals
1A2d	Stationary combustion in manufacturing industries and construction: Pulp, Paper and Print
1A2e	Stationary combustion in manufacturing industries and construction: Food processing, beverages and tobacco
1A2f	Stationary combustion in manufacturing industries and construction: Non-metallic minerals
1A2gviii	Stationary combustion in manufacturing industries and construction: Other

This chapter presents the consumption of fuels and emissions of air pollutants in six specific types of industry, all other industries are hidden under NFR Code 1A2gviii, Stationary combustion in manufacturing industries and construction: Other. NFR Code 1A2gviii includes a big number of enterprises. In addition, fuel for construction is included under 1A2gviii: Other, except diesel and gasoline. Diesel and gasoline are included under 1A2gvii: Mobile Combustion in manufacturing industries and construction.

#### Methodology

To estimate emissions from combustion in manufacturing industries and construction the following formulas have been used:

$$E = m \times \text{NCV} \times \text{EF} \quad \text{Equation 1}$$

E - emission (g)  
m - quantity of fuel combusted (t)  
NCV - net calorific value (TJ/kt)  
EF - emission factor per energy of fuel (g/GJ)

$$E = m \times \text{EF} \quad \text{Equation 2}$$

E - emission (g)  
m - quantity of fuel combusted (t)  
EF - emission factor per quantity of fuel (g/t)

To estimate SO<sub>x</sub> emissions in some cases the following two equations for calculation of EF were used:

$$EF_{SO_x} = [S] \times 20000 / NCV \quad \text{Equation 3}$$

$EF_{SO_x}$  –  $SO_x$  emission factor (g/GJ)  
 [S] – sulphur content of the fuel (% w/w)  
 NCV - net calorific value (GJ/t)  
 2 – ratio of the relative molecular mass of  $SO_2$  to sulphur

$$EF_{SO_x} = [S] \times 19000 / NCV \quad \text{Equation 4}$$

$EF_{SO_x}$  -  $SO_x$  emission factor (g/GJ)  
 [S] – sulphur content of the fuel (% w/w)  
 NCV - net calorific value (GJ/t)  
 1.9 – ratio of the relative molecular mass of  $SO_2$  to sulphur, considering 5 % absorption in the ash

The total emission for this sub/sector is the sum of different industrial activities, using diverse fuels and combustion technologies.

### Activity data

The fuel consumption in each category has to be determined in accordance with the classification of activities applied in EMEP/EEA emission inventory guidebook, 2013.

#### Period 1980-1996

**Table 3.2.1.1 Conversion table between national energy statistics (LEG) and NFR category**

NFR category	LEG Classification (1986-1996)
Iron and Steel	Iron and Steel Production
Non-Ferrous Metals	Non-Ferrous Metals
Chemicals	Chemical Industry
Pulp, Paper and Print	Pulp and Paper Industry, Print Industry
Food Processing, Beverages and Tobacco	Food Processing Industry, Tobacco Industry
Non-metallic minerals	Non-metal industry
Other	Metal Industry Shipbuilding Electrical Industry Construction Timber Industry Textile Industry Leather Industry Rubber Industry Recycling Other Industry

The classification applied in LEG has been taken as the basis and conversion table between LEG and NFR is presented in the Table 3.2.1.1.

#### Period 1997-2003

In 1997, LEG began to publish data according to the Standard Classification of Activities (SCA) which in some categories differs from the classification, which had been used until 1996. Most activities are defined in a similar manner, but this is not possible for certain activities. The table 3.2.1.2 shows the distribution of activities in accordance with the EMEP/EEA classification.

For consumption in individual industrial sectors there are detailed (disaggregated) data, the values of which was strongly dependant on the mode of reporting and features of individual

industrial sectors characterized by high concentration (values depending on the consumption in one or two factories) in Slovenia. Data from basic sources hint at some relatively big changes in the consumption of fuels in some sectors.

**Table 3.2.1.2 Conversion table between national energy statistics (LEG) and NFR**

NFR category	LEG Classification – SCA category
Iron and Steel	DJ - Production of metals and metal products
Non-Ferrous Metals	
Chemicals	DG - Production of chemicals
Pulp, Paper and Print	DE - Production of fibres, pulp, paper, and cardboard
Food Processing, Beverages and Tobacco	DA – Production of food, beverages, and tobacco products
Non-metallic Minerals	DI - Production of non-metal mineral products
Other	DB - Production of textiles
	DC - Production of leather and leather goods
	DD – Wood-processing and woodworking
	DH - Production of rubber products
	DK - Production of machines and devices
	DL - Production of electrical and optical equipment
	DM – Production of vehicles and vessels
	DN - Production of furniture. not included elsewhere
F - Construction	

#### Period 2004 - 2007

Since 2004 very detailed data about fuel consumption in industry become available in electronic format. The non-energy and energy use of fuels are reported separately. Data about fuel consumption and NCV are reported on the lowest level of disaggregation possible. For this reason, from 2004 on fuel consumption in iron and steel industry and in non-ferrous metals industry can be separated according to the rules presented in the following Table 3.2.1.3.

**Table 3.2.1.3 Table for disaggregation of fuel in DJ sector (manufacture of basic metals and fabricated metal products)**

SCA category	NFR category	Description
DJ 27.1	Iron and Steel	Manufacture of basic iron and steel and of ferrous alloys
DJ 27.2	Iron and Steel	Manufacture of tubes
DJ 27.3	Iron and Steel	Other first processing of iron and steel
DJ 27.4	Non-ferrous Metal	Manufacture of basic precious and non-ferrous metals
DJ 27.510	Iron and Steel	Casting of iron
DJ 27.520	Iron and Steel	Casting of steel
DJ 27.530	Non-ferrous Metal	Casting of light metal
DJ 27.540	Non-ferrous Metal	Casting of other non-ferrous metal
DJ 28	Other industry	Manufacture of fabricated metal products, except machinery and equipment

#### Period 2008 – 2017

**Table 3.2.1.4 Conversion table between the NFR categories and The Standard Classification of Activities (SKD)**

NFR category	Description
1.A.2.a Iron and Steel	C 24.1 Manufacture of basic iron and steel and of ferrous alloys
	C 24.2 Manufacture of tubes, pipes, hollow profiles and related fittings, of steel
	C 24.3 Manufacture of other products of first processing of steel
	C 24.51 Casting of iron

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	C 24.52 Casting of steel
1.A.2.b Non-ferrous Metal	C 24.4 Manufacture of basic precious and non-ferrous metals
	C 24.53 Casting of light metal
	C 24.54 Casting of other non-ferrous metal
1.A.2.c Chemicals	C 20 Manufacture of chemicals and chemical products
1.A.2.d Pulp, Paper and Print	C 17 Manufacture of paper and paper products
	C 18 Printing and reproduction of recorded media
1.A.2.e Food Processing, Beverages and Tobacco	C 10 Manufacture of food products
	C 11 Manufacture of beverages
	C 12 Manufacture of tobacco products
1.A.2.f Non-metallic Minerals	C 23 Manufacture of other non-metallic mineral products
1.A.2.g.vii Off road vehicles and other machinery	F Construction (only gasoline and diesel fuel)
1.A.2.g.viii Other	C 13 Manufacture of textiles
	C 14 Manufacture of wearing apparel
	C 15 Manufacture of leather and related products
	C 16 Manufacture of wood and of products of wood and cork, except furniture, manufacture of articles of straw and plaiting materials
	C 21 Manufacture of basic pharmaceutical products and pharmaceutical preparations
	C 22 Manufacture of rubber and plastic products
	C 25 Manufacture of metallic products
	C 26 Production of electrical and optical equipment
	C 27 Production of electrical equipment
	C 28 Production of machines and devices
	C 29 Production of vehicles
	C 30 Production of vessels
	C 31 Production of furniture
	C 32 Other manufacturing
	C 33 Repair and installation of machinery and equipment
F Construction (all other fuels except diesel and gasoline)	

In 2008 the new SCA (Standard Classification of Activities) was applied by SORS which was used until present. The main advantage is that the new classification enables disaggregation of data on much more detailed level. An important difference is that "Manufacture of basic pharmaceutical products and pharmaceutical preparations" industry is no longer part of the Chemical industry and is included under category "Other". The conversion table between NFR and national energy statistics is presented in the Table 3.2.1.4.

In industry, particularly in cement industry, in addition to commonly used fuel, some waste is also incinerated because of very high temperature in the oven. We have obtained very detailed data about amount and composition of waste from one cement plant, where the main process of waste incineration in Slovenia was occurring. Since 2005, all waste fuels have also been included in ETS.

We had also obtained data from pulp and paper industry about consumption of black liquor from 2004 to 2006. NCV was between 6,1 and 6,4 TJ/kt. We used the same emissions factors for calculation as for wood. From 2007, there has been no consumption of black liquor any more.

## **Inclusion of auto producers into Manufacturing Industries sector**

In accordance with IPCC Reference manual, the item Industry reports the consumption of fuels in the group of industrial power plants (auto producers – enterprises that generate electric energy for internal consumption and/or heat for sale) as well as other consumption in industry (except in production processes). The same methodology was adopted also for emission calculation of air pollutants.

In the period 1986 -1996, consumption of fuels by auto producers in LEG was recorded under Electric utilities – Industry, and in the period 1997- 2003 under Conversion – Auto producers.

### Period 1986-2000

Because there are no published data on auto producers at the level of industrial branches for the period 1986-2000, on the basis of which it would be possible to assign the consumption of fuel to each individual industrial branch, for each kind of fuel a different (most appropriate) approach was used.

#### Lignite

Total consumption is attributed to pulp and paper industry. The paper mill in Krško uses lignite in its power cogeneration plant. In the documents of the SORS, the total consumption is attributed to the consumption in thermal power plants, while in LEG one half of the consumption is attributed to the consumption in industry, the other half to industrial thermal power plants. In this report, a half is reported as consumption in pulp and paper industry (heat), a half as consumption in industrial power plants in pulp and paper industry. Consumption of lignite in other sectors has not been reported.

#### Brown Coal

Consumption of brown coal in industrial power plants in the monitored period was reported only in 1986. Since quantities are quite small, consumption is reported in the sector “Other”.

#### Residual Fuel Oil

Consumption of residual fuel oil in industrial power plants in the monitored period was low (from 0 to 10176 t). Since quantities are quite small, consumption is reported in sector “Other”.

#### Gas Oil and Natural Gas

The majority of industrial thermal power plants use gas oil or natural gas. Total quantities of consumed gas oil and natural gas are disaggregated according to the produced quantities of electric energy in those power plants.

### Period 2000-2017

Since 2000 we have commenced to treat auto producers individually, since the SORS, which prepares data for LEG, has completed its database. Now, aggregated data on the consumption of fuels by auto producers at the level of industrial branches are available, where the sums of individual fuels correspond to the consumption of auto producers from LEG.

Following the recommendations of the expert review team data on fuel consumption by industry type, fuel type and year are reported in the Annex 1 to the IIR, (Table 1.15: Fuel used in Manufacturing industries and construction 1980–2017).

### Net calorific values

Tables 3.2.1.5 to 3.2.1.8 present the net calorific values (NCV) which have been used for fuel combusted in manufacturing industries. In the past they have been mostly taken from Statistical Office of the Republic of Slovenia while since 2005 the ETS data are used, if available. Plant specific data for 2017 for solid fuels are presented in the Table 3.2.1.7. The values for liquid fuels excluding petrol coke, natural gas and biomass have been taken from SORS for the entire period.

**Table 3.2.1.5 NCVs for the fuel used in manufacturing industry and construction**

Year	Lignite – domestic (Velenje)	Sub-bituminous Coal - domestic	Lignite - imported	Sub-bituminous Coal - imported	Other Bituminous Coal	Anthracite	Coke	Petroleum coke
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt
1980	9,390	11,880			27,57	29,25	29,30	31,000
1981	9,390	11,880			27,57	29,25	29,30	31,000
1982	9,390	11,880			27,57	29,25	29,30	31,000
1983	9,390	11,880			27,57	29,25	29,30	31,000
1984	9,650	11,820			27,57	29,25	29,30	31,000
1985	9,390	11,880			27,57	29,25	29,30	31,000
1986	9,390	11,880			27,57	29,25	29,30	31,000
1987	9,650	11,820			27,57	29,25	29,30	31,000
1988	9,440	12,000			27,57	29,25	29,30	31,000
1989	9,820	12,050			27,57	29,25	29,30	31,000
1990	9,810	12,760			27,57	29,25	29,30	31,000
1991	9,980	12,879			25,00	29,25	29,30	31,000
1992	10,260	12,589			25,00	29,25	29,30	31,000
1993	10,070	13,351			25,00	29,25	29,30	31,000
1994	9,960	12,666			25,00	29,25	29,30	31,000
1995	10,220			17,404	25,00	29,31	29,31	31,000
1996	9,690			16,353	25,00	29,31	29,31	31,000
1997	9,610			17,712	25,00	29,31	29,310	31,000
1998	10,010			20,664	25,00	29,31	29,310	31,000
1999	9,690			20,806	25,00	29,31	29,310	31,000
2000	10,170			20,782	25,00	29,31	29,310	31,000
2001	10,660			20,947	25,00	29,31	29,310	31,000
2002	10,350			21,000	25,00	29,31	29,310	31,000
2003	10,138			21,570	25,00	29,31	29,310	31,000
2004	10,301			19,908		29,40	30,031	29,927

**Table 3.2.1.6 NCVs for the fuel used in manufacturing industry and construction**

Year	Residual Fuel Oil	Heavy Fuel Oil	Diesel	Gasoline	LPG	Natural Gas
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/Mm3
1980	41,82	39,74	42,70	43,18	46,00	33,50
1981	41,82	39,74	42,70	43,18	46,00	33,50
1982	41,82	39,74	42,70	43,18	46,00	33,50
1983	41,82	39,74	42,70	43,18	46,00	33,50
1984	41,82	39,74	42,70	43,18	46,00	33,50
1985	41,82	39,74	42,70	43,18	46,00	33,50
1986	41,82	39,74	42,70	43,18	46,00	33,50
1987	41,78	39,80	42,70	43,10	46,00	33,50
1988	41,71	39,80	42,70	43,10	46,00	34,08

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1989	41,85	39,80	42,70	43,10	46,00	34,10
1990	41,87	39,80	42,70	43,07	46,00	34,10
1991	41,88	39,80	42,70	43,17	46,00	34,10
1992	41,90	39,90	42,70	43,10	46,00	34,10
1993	41,90	39,80	42,70	43,08	46,00	34,10
1994	41,90	39,86	42,70	43,08	46,00	34,10
1995	41,90	40,00	42,70	43,08	46,00	34,10
1996	41,90	40,00	42,70	43,08	46,00	34,10
1997	41,90	40,00	42,70	43,08	46,05	34,08
1998	41,90	40,00	42,70	43,08	46,05	34,08
1999	41,90	40,00	42,70	43,08	46,05	34,08
2000	41,90	40,00	42,70	43,08	46,05	34,08
2001	41,90	40,00	42,70	43,08	46,05	34,08
2002	41,90	40,00	42,70	43,08	46,05	34,08
2003	41,90	40,00	42,70	43,08	46,05	34,08
2004	41,90	40,00	42,70	43,08	46,05	34,08
2005	42,60	41,42	42,70	43,08	46,05	34,08
2006	42,60	41,42	42,70	43,08	46,05	34,07
2007	42,60	41,42	42,70	43,08	46,05	34,08
2008	42,60	41,42	42,70	43,85	46,05	34,09
2009	42,60	41,42	42,70	43,85	46,05	34,08
2010	42,60	41,42	42,70	43,85	46,05	34,08
2011	42,60	41,42	42,60	43,85	46,05	34,09
2012	42,60	41,42	42,60	43,85	46,05	34,09
2013	42,60	41,42	42,60	43,85	46,05	34,08
2014	42,60	41,42	42,60	43,85	46,05	34,08
2015	42,60	41,42	42,60	43,85	46,05	34,08
2016	42,60	41,42	42,60	43,85	46,05	34,08
2017	42,60	41,42	42,60	43,85	46,05	34,08

Table 3.2.1.7 NCVs for the solid fuel used in manufacturing industry and construction in 2017

Industry	Unit	Sub-bituminous Coal - imported	Other Bituminous Coal	Coke	Petroleum coke	Wood	Other biomass
Iron and steel	TJ/kt			30,166		15,500	
Non-Ferrous metals	TJ/kt		25,000				
Chemicals	TJ/kt					10,680	
Pulp. Paper and Print	TJ/kt	19,473				8,863	3,481
Food processing	TJ/kt					12,621	
Non-metallic minerals	TJ/kt			29,300	31,263	12,601	13,789
Other	TJ/kt	21,000				12,575	

Table 3.2.1.8 NCVs for other fuels

	Waste industrial oils	Waste cooking fat	Waste cooking oils	Waste tyres	Waste organic solvents	Other waste
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt
1996	37,00			27,21		11,00
1997	37,00			27,21		11,00
1998	37,00			27,21		11,00



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1999	37,00			27,21		11,00
2000	37,00			27,21		11,00
2001	37,00	39,20		27,21		11,00
2002	37,00	39,20		27,21		11,00
2003	37,00	39,20		27,21		11,00
2004	41,90	40,41	40,00	27,21		
2005	34,64	39,20	40,00	27,21		
2006	34,53	39,20		27,21		
2007	33,76	39,95		27,21		
2008	34,48	39,81		27,21		17,52
2009	37,65	39,81		27,19		26,67
2010	36,95	39,20		27,23	25,00	22,34
2011	36,25	39,20		27,26	25,00	19,52
2012	37,09	39,20		27,21	25,00	20,25
2013	37,13	39,20		27,21	25,00	19,44
2014	33,03	39,20		27,20	25,00	18,87
2015	35,49	39,20		27,20	25,00	19,32
2016	36,54	39,20		27,20	25,00	18,19
2017	37,34	38,23		27,20	25,00	16,90

### Emission factors

For calculating emissions of individual gases in manufacturing industry and construction following emission factors have been used.

**Table 3.2.1.9 Emission factors used for domestic sub-bituminous coal, imported sub-bituminous coal, domestic and imported lignite, other bituminous coal, anthracite and coke for 1980 - 2017**

Pollutant	Value	Unit	References
NO <sub>x</sub>	173	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
SO <sub>x</sub>	Equation 4	[S] (% w/w) See Table 3.2.1.10	Slovene national legislation relating quality of liquid fuels
CO	931	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
NM VOC	88,8	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
PM <sub>10</sub>	117	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
PM <sub>2.5</sub>	108	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
TSP	124	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
BC	6,91	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
Cd	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
Pb	134	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
Hg	7,9	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
As	4	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
Cr	13,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15

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<b>Cu</b>	17,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Ni</b>	13	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Se</b>	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Zn</b>	200	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Dioxins/ Furans</b>	203	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Benzo(a)pyrene</b>	45,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Benzo(b)fluoranthene</b>	58,9	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Benzo(k)fluoranthene</b>	23,7	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>Indeno(1,2,3-cd)pyrene</b>	18,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>HCB</b>	0,62	microg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15
<b>PCB</b>	170	microg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-2, pg 15

Table 3.2.1.10 Sulphur content in coals, anthracite and coke for 1980 - 2017

Year	Lignite domestic/imported	Sub-bituminous Coal - domestic	Sub-bituminous Coal - imported	Other Bituminous Coal	Anthracite	Coke/ Petroleum coke
	[S] (% w/w)	[S] (% w/w)	[S] (% w/w)	[S] (% w/w)	[S] (% w/w)	[S] (% w/w)
1980	1,45	1,6		8	1	1
1981	1,45	1,6		8	1	1
1982	1,45	1,6		8	1	1
1983	1,45	1,6		8	1	1
1984	1,45	1,6		8	1	1
1985	1,45	1,6		8	1	1
1986	1,45	1,6		8	1	1
1987	1,45	1,6		8	1	1
1988	1,45	1,6		8	1	1
1989	1,45	1,6		8	1	1
1990	1,45	1,6		8	1	1
1991	1,45	1,6		8	1	1
1992	1,45	1,6		8	1	1
1993	1,45	1,6		8	1	1
1994	1,45	1,6		8	1	1
1995	1,45		1,60	8	1	1
1996	1,45		1,60	8	1	1
1997	1,45		1,60	8	1	1
1998	1,45		0,12	8	1	1
1999	1,45		0,12	8	1	1
2000	1,45		0,12	8	1	1
2001	1,45		0,12	8	1	1
2002	1,45		0,07	1	1	1
2003	1,45		0,09	1	1	1
2004	1,45		0,09	1	1	1
2005			0,14	1		1
2006			0,14	1		1
2007			0,14	1		1
2008			0,10	1		1

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2009	1,45		0,10	1		1
2010	1,45		0,10	1		1
2011	1,45		0,10	1		1
2012	1,45		0,10	1		1
2013	1,45		0,10	1		1
2014	1,45		0,10	1		1
2015	1,45		0,10	1		1
2016	1,45		0,10	1		1
2017			0,10	1		1

Table 3.2.1.11 Emission factors used for heavy fuel, residual fuel oil, petroleum coke, waste industrial oils and waste organic solvents for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	513	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
SO <sub>x</sub>	Equation 3	[S] (% w/w) See Table 3.2.1.12	Slovene national legislation relating quality of liquid fuels
CO	66	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
NM VOC	25	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
PM <sub>10</sub>	20	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
PM <sub>2.5</sub>	20	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
TSP	20	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
BC	11,2	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Cd	0,006	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Pb	0,08	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Hg	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
As	0,03	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Cr	0,20	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Cu	0,22	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Ni	0,008	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Se	0,11	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Zn	29	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Benzo(a)pyrene	1,9	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Benzo(b)fluoranthene	15	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Benzo(k)fluoranthene	1,7	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Indeno(1,2,3-cd)pyrene	1,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17
Dioxins/ Furans	1,4	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-4, pg 17

Table 3.2.1.12 Sulphur content in residual fuel oil and heavy fuel oil for 1980 – 2017

Fuel	Heavy fuel Oil	Residual fuel Oil	Fuel	Heavy fuel Oil	Residual fuel Oil
Year	[S] (% w/w)	[S] (% w/w)	year	[S] (% w/w)	[S] (% w/w)
1980	3,0	1,2	1999	1,0	0,2
1981	3,0	1,2	2000	1,0	0,2
1982	3,0	1,2	2001	1,0	0,2
1983	3,0	1,2	2002	1,0	0,2
1984	3,0	1,2	2003	1,0	0,2
1985	3,0	1,2	2004	1,0	0,2
1986	3,0	1,2	2005	1,0	0,2
1987	3,0	1,2	2006	1,0	0,2
1988	3,0	1,2	2007	1,0	0,2
1989	3,0	1,2	2008	1,0	0,1
1990	3,0	1,2	2009	1,0	0,1
1991	3,0	1,2	2010	1,0	0,1
1992	3,0	1,2	2011	1,0	0,1
1993	3,0	1,2	2012	1,0	0,1
1994	3,0	1,2	2013	1,0	0,1
1995	1,5	0,5	2014	1,0	0,1
1996	1,0	0,2	2015	1,0	0,1
1997	1,0	0,2	2016	1,0	0,1
1998	1,0	0,2	2017	1,0	0,1

Table 3.2.1.13 Emission factors used for wood, other biomass, waste cooking fat and waste cooking oils for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	91	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
CO	570	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
NM VOC	300	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
SO <sub>x</sub>	11	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
NH <sub>3</sub>	37	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
PM <sub>10</sub>	143	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
PM <sub>2.5</sub>	140	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
TSP	150	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
BC	39,2	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Cd	13	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Pb	27	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Hg	0,56	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
As	0,19	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18

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Cr	23	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Cu	6	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Ni	2	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Se	0,5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Zn	512	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Benzo(a)pyrene	10	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Benzo(b)fluoranthene	16	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Benzo(k)fluoranthene	5	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Indeno(1,2,3-cd)pyrene	4	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
Dioxins/ Furans	100	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
PCB	0,06	microg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18
HCB	5	microg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-5, pg 18

Table 3.2.1.14 Emission factors used for natural gas, biogas and liquefied petroleum gas for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	74	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
CO	29	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
SO <sub>x</sub>	0,67	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
NMVOC	23	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
PM <sub>10</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
PM <sub>2.5</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
TSP	0,78	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
BC	0,0312	g/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Cd	0,0009	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Pb	0,011	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Hg	0,54	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
As	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Cr	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Cu	0,0026	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Ni	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Se	0,058	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
Zn	0,73	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16

<b>Benzo(a)pyrene</b>	0,72	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
<b>Benzo(b)fluoranthene</b>	2,9	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
<b>Benzo(k)fluoranthene</b>	1,1	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
<b>Indeno(1,2,3-cd)pyrene</b>	1,08	mg/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16
<b>Dioxins/ Furans</b>	0,52	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Manufacturing industries and Construction, Table 3-3, pg 16

Table 3.2.1.15 Emission factors used for waste tyres and other waste

Pollutant	Value	Unit	References
<b>NO<sub>x</sub></b>	0,87	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>SO<sub>x</sub></b>	0,047	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>CO</b>	0,07	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>NMVOG</b>	7,4	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>PM<sub>2.5</sub></b>	0,004	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>PM<sub>10</sub></b>	0,007	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>TSP</b>	0,01	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>BC</b>	0,00014	kg/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>Cd</b>	0,1	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>Hg</b>	0,056	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>Pb</b>	1,3	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>As</b>	0,016	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>Ni</b>	0,14	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>Dioxins/ Furans</b>	1	microg I-TEQ/t	Plant specific
<b>Total 4 PAHs</b>	0,02	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1
<b>HCB</b>	0,002	g/t	Emission Inventory Guidebook, 2016, Industrial waste incineration, pg 10, Table 3-1

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors for biomass represent filterable PM emissions. It is unclear for solid fuels, gaseous fuel, liquid fuels and waste whether emission factors represent filterable PM emissions or total (filterable and condensable) emissions.

## Emissions

Manufacturing industries and construction sector is significant source of emissions. In 2017 contributed about 16 % to total national SO<sub>x</sub> emissions, 10 % to NO<sub>x</sub>, 8 % to NMVOG, 7 % to particulate, 18 % to Hg. Emissions of almost all pollutants have declined in the last decades due to improvement in technologies, implementation of abatement techniques and fuel switching to cleaner fuels.

## **Recalculations**

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Due to new activity data on natural gas obtained for 2016 in Other manufacturing industries and construction recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene and indeno(1,2,3-cd)pyrene was performed for the year 2016.

Activity data on anthracite was obtained for 1996 in Other manufacturing industries and construction sub sector. Recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, indeno(1,2,3-cd)pyrene, HCB and PCB was performed for the year 1996.

## **Category-specific QA/QC and verification**

The source category QA/QC is covered by the general QC procedures described in the chapter 2.5. Our main source specific QA/QC activity is comparison of the ETS data with statistical data. The aggregated fuel from SORS data is compared with the sum of fuel used from verified ETS reports and where connection between both set of data is uniform, the data from SORS are substituted with data from the verified reports from installations included in ETS, if necessary. ETS data are also used for different types of waste used as fuel. The list of waste types is not always complete in the SORS data. Additional QA activity is reference approach. Before entering data into database, the sum of each fuel from disaggregated data is compared with energy balance data, reported in the Joint Questioner. As data in JQ are rounded to 1000 units, the difference should be 500 units or less. If it is higher, the reasons for this should be found.

## **Future improvements**

No improvements are planned for next submission.

### 3.3 Transport (1. A. 3)

Transport is an important source of emissions of air pollutants, mostly nitrous oxide. It is also an important source of other emissions what cause problems in terms of air quality. The most important source in category transport is road transport, which accounts more than 95 % of all transport emissions.

Sectors covered in this chapter are:

NFR Codes:

1A3bi -1A3bvii	Road transport
1A3c	Railways
1A3ai(i)	International aviation LTO (civil)
1A3aii(i)	Domestic aviation LTO (civil)
1A5b	Other, Mobile (including military, land based and recreational boats)

Emissions from sectors NFR Code 1A5c Multilateral operations and NFR Code 1A3di(i) International maritime navigation are reported under Memo items. Emissions are therefore not included in national total emissions.

#### 3.3.1 Road transport

Sectors covered in this chapter are:

NFR Codes:

1A3bi	Road transport: Passenger cars
1A3bii	Road transport: Light duty vehicles
1A3biii	Road transport: Heavy duty vehicles and buses
1A3biv	Road transport: Mopeds & motorcycles
1A3bv	Road transport: Gasoline evaporation
1A3bvi	Road transport: Automobile tyre and brake wear
1A3bvii	Road transport: Automobile road abrasion

#### Introduction

Road transportation is one of the most important emitter of greenhouse gases (GHG) such as carbon dioxide (CO<sub>2</sub>), methane (CH<sub>4</sub>) and nitrous oxide (N<sub>2</sub>O). It is also a significant emission source of pollutants associated with trans-boundary, regional and local air problems, comprehending sulphur oxides (SO<sub>x</sub>), nitrogen oxides (NO<sub>x</sub>), carbon monoxide (CO), non-volatile organic compounds (NMVOC) and are indirectly responsible for the formation of ozone (O<sub>3</sub>) in the lower troposphere. Substantial emissions of ammonia (NH<sub>3</sub>), particulate matter (PM) and heavy metals also result from this activity.

#### Methodology

COPERT 4 (version 11.4) methodology has been used for the calculation of national emission estimates from road transport for the entire 1980-2017 period. The methodology is fully incorporated in the computer software programme COPERT 4 which facilitates its application. The actual calculations have been therefore performed by using this computer software.



COPERT 4 estimates emissions of all major air pollutants (CO, NO<sub>x</sub>, NMVOC, particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, Black carbon), NH<sub>3</sub>, SO<sub>x</sub>, heavy metals) as well as greenhouse gas emissions (CO<sub>2</sub>, N<sub>2</sub>O, CH<sub>4</sub>) produced by different vehicle categories (passenger cars, light duty vehicles, heavy duty trucks, buses, mopeds and motorcycles). The programme also provides speciation of polyaromatic hydrocarbons (PAHs) and dioxins/furans. Emissions of HCB and PCB are given as a total emissions from road transport. Emissions estimated are distinguished in three sources: emissions produced during thermally stabilized engine operation (hot emissions), emissions occurring during engine start from ambient temperature (cold-start and warming-up effects) and NMVOC emissions due to fuel evaporation. The total emissions are calculated as a product of activity data provided by the user and speed-dependent emission factors calculated by the software.

The COPERT 4 methodology is also part of the EMEP/EEA air pollutant emission inventory guidebook (formerly referred to as the EMEP/ CORINAIR Guidebook). The Guidebook is prepared by the UNECE/EMEP Task Force on Emission Inventories and Projections (TFEIP) and published by the European Environment Agency. It is intended to support reporting under the UNECE Convention on Long-Range Transboundary Air Pollution and the EU directive on national emission ceilings as well as under United Nations Framework Convention on Climate Change (UNFCCC). The COPERT 4 methodology is fully consistent with the Road Transport chapter of the Guidebook. The use of a software tool to calculate road transport emissions allows for a transparent and standardized, hence consistent and comparable data collecting and emissions reporting procedure, in accordance with the requirements of international conventions and protocols and EU legislation.

Applied methodology is fully described in the following literature:

- COPERT 4 Computer programme to calculate emissions from road transport - User manual (version 5.0), Dimitrios Gkatzoflias, Chariton Kouridis, Leonidas Ntziachristos and Zissis
- EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Chapters: 1.A.3.b.i-iv Exhaust emissions from road transport, 1.A.3.b.v Gasoline evaporation.

To calculate emissions using the COPERT 4 software, at least the following input data is necessary: vehicle fleet data, mileage data per vehicle category and type of roads, speed data, fuel consumption and fuel characteristic, monthly air minimum and maximum temperatures, fuel vapour pressure.

COPERT 4 (version 11.4) programme was concretely used for emissions calculation of NO<sub>x</sub>, SO<sub>x</sub>, NMVOC, NH<sub>3</sub>, PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, Black carbon (BC), CO, Lead (Pb), Cadmium (Cd), Chromium (Cr), Copper (Cu), Nickel (Ni), Selenium (Se), Zinc (Zn), dioxins/furans and four indicator PAHs (benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, indeno(1,2,3-cd)pyrene), PCB and HCB.

Emissions of particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC) from automobile tyre and brake wear and road abrasion have been calculated using methodology and emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Chapters: 1.A.3.b.vi and 1.A.3.b.vii. Road transport: Automobile tyre and brake wear, Automobile road abrasion. Emissions of heavy metals (Pb, Cd, Cu, Cr, Ni Se, Zn) from automobile tyre and brake wear have been calculated using COPERT 4 (version 11.4).

## Vehicle fleet

The COPERT 4 methodology requires a detailed knowledge of the structure of the vehicle fleet composition Table 3.3.1.1 provides a summary of all vehicle categories and technologies covered by the applied methodology.

The fleet composition for the years 1992–2009 was taken from the official database of registered motor and trailer vehicles in the Republic of Slovenia provided by the Ministry of the Interior. Since 2010 these data have been collected by Ministry of Infrastructure of the Republic of Slovenia. Since no database exists on licensed motor and trailer vehicles in the Republic of Slovenia for the years 1980–1991, an expert estimate has been made on the basis of the annual Statistical Yearbooks, published by Statistical Office of the Republic of Slovenia (SORS).

The vehicle numbers per all vehicle classes for period 1980–2017 are shown in the Annex 1 to the IIR (Table 1.1 Road transport: Fleet data (number of vehicles) 1980–2017).

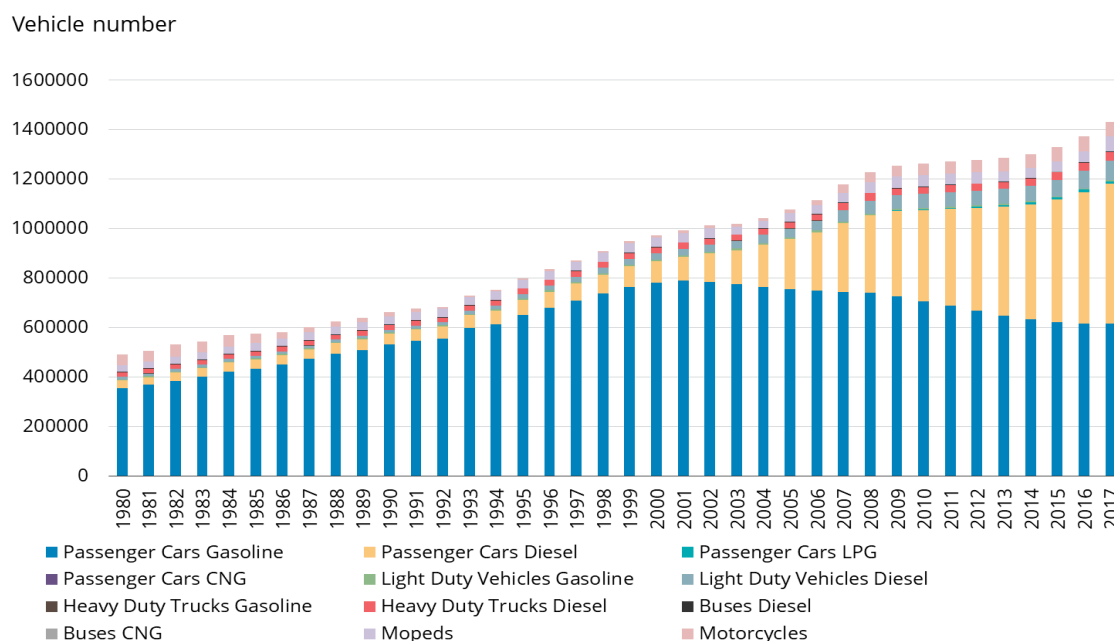
**Table 3.3.1.1 Summary of vehicle classes covered by the methodology**

Vehicle Type	Class	Legislation
Passenger Cars	Gasoline <1.4l	PRE ECE ECE 15/00-01 ECE 15/02 ECE 15/03 ECE 15/04
	Gasoline 1.4 - 2.0l	Improved Conventional Open Loop Euro 1 - 91/441/EEC Euro 2 - 94/12/EEC
	Gasoline >2.0l	Euro 3 - 98/69/EC Stage 2000 Euro 4 - 98/69/EC Stage 2005 Euro 5 - EC 715/2007 Euro 6 - EC 715/2007 Euro 6c - EC 715/2007
	Diesel <2.0l	Conventional Euro 1 - 91/441/EEC Euro 2 - 94/12/EEC
	Diesel >2.0l	Euro 3 - 98/69/EC Stage 2000 Euro 4 - 98/69/EC Stage 2005 Euro 5 - EC 715/2007 Euro 6 - EC 715/2007 Euro 6c - EC 715/2007
	LPG	Conventional Euro 1 - 91/441/EEC Euro 2 - 94/12/EC Euro 3 - 98/69/EC Stage 2000 Euro 4 - 98/69/EC Stage 2005 Euro 5 — EC 715/2007 Euro 6 — EC 715/2007
	2 Stroke	Conventional
	Hybrids Gasoline <1.4l Hybrids Gasoline 1.4-2.0l Hybrid Gasoline >2.0l	Euro 4 - 98/69/EC Stage 2005
Light Duty Vehicles	Gasoline <3.5t	Conventional Euro 1 - 93/59/EEC Euro 2 - 96/69/EEC Euro 3 - 98/69/EC Stage 2000 Euro 4 - 98/69/EC Stage 2005 Euro 5 - EC 715/2007 Euro 6 - EC 715/2007 Euro 6c - EC 715/2007

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	Diesel <3.5t	Conventional Euro 1 - 93/59/EEC Euro 2 - 96/69/EC Euro 3 - 98/69/EC Stage 2000 Euro 4 - 98/69/EC Stage 2005 Euro 5 - EC 715/2007 Euro 6 - EC 715/2007 Euro 6c - EC 715/2007
Heavy Duty Trucks	Gasoline >3.5t	Conventional
	Rigid <=7.5t	Conventional Euro I - 91/542/EEC Stage I Euro II - 91/542/EEC Stage II Euro III - 1999/96/EC Stage I Euro IV - 1999/96/EC Stage II Euro V - 1999/96/EC Stage III Euro VI - Regulation EC 595/2009
	Rigid 7.5-12t	
	Rigid 12-14t	
	Rigid 14-20t	
	Rigid 20-26t	
	Rigid 26-28t	
	Rigid 28-32t	
	Rigid >32t	
	Articulated 14-20t	
	Articulated 20-28t	
	Articulated 28-34t	
	Articulated 34-40t	
	Articulated 40-50t	
Articulated 50-60t		
Buses	Urban <=15t	Conventional Euro I - 91/542/EEC Stage I Euro II - 91/542/EEC Stage II Euro III - 1999/96/EC Stage I Euro IV - 1999/96/EC Stage II Euro V - 1999/96/EC Stage III Euro VI - Regulation EC 595/2009
	Urban 15-18t	
	Urban >18t	
	Coaches articulated >18t	
	Coaches standard <=18t	
	CNG	
Mopeds	2-stroke, < 50 cm <sup>3</sup>	Conventional Euro 1 - 97/24/EC Stage I Euro 2 - 97/24/EC Stage II Euro 3 - Directive 2002/51/EC Euro 4 - Regulation EC 168/2013 Euro 5 - Regulation EC 168/2013
	4-stroke, < 50 cm <sup>3</sup>	
Motorcycles	2-stroke, > 50 cm <sup>3</sup>	Conventional 97/24/EC – Euro 1 2002/51/EC Stage I - Euro 2 2002/51/EC Stage II - Euro 3 Euro 4 - Regulation EC 168/2013 Euro 5 - Regulation EC 168/2013
	4-stroke, 50–250 cm <sup>3</sup>	
	4-stroke, 250–750 cm <sup>3</sup>	
	4-stroke, > 750 cm <sup>3</sup>	

The vehicle fleet structure is presented in Figure 3.3.1.1. The increase in the total number of passenger cars is mostly due to a growth in the number of diesel passenger cars. After the year 2003 a considerable decline in the number of gasoline passenger cars is observed, and at the same time a rise in the number of diesel passenger cars. LPG and CNG passenger cars represent only a small share of all passenger cars



**Figure 3.3.1.1 Vehicle fleet 1980–2017**

### Mileage

Annual mileage (km/year) for each vehicle category have been obtained from the Ministry of Infrastructure of the Republic of Slovenia, SORS and official database of registered motor and trailer vehicles in the Republic of Slovenia provided by the Ministry of Infrastructure of the Republic of Slovenia. The values used are shown in the Annex 1 (Table 1.2: Road transport: Mileage data 1980–2017).

### Mileage driven and number of vehicles for particulates from tyre and brake wear and road abrasion

The activity data, vehicle kilometres per vehicle category and number of vehicle per vehicle category, needed for calculation particulate matter ( $PM_{2.5}$ ,  $PM_{10}$ , TSP, BC) emissions from automobile tyre and brake wear and road abrasion have been derived from Copert 4 model, version 11.4. Source of original data (mileage and vehicle fleet) are presented in previous paragraphs of this chapter. The values used are shown in the Annex 1 (Table 1.3: Road transport: particulates from tyre and brake wear and road abrasion 2000–2017).

### Speed

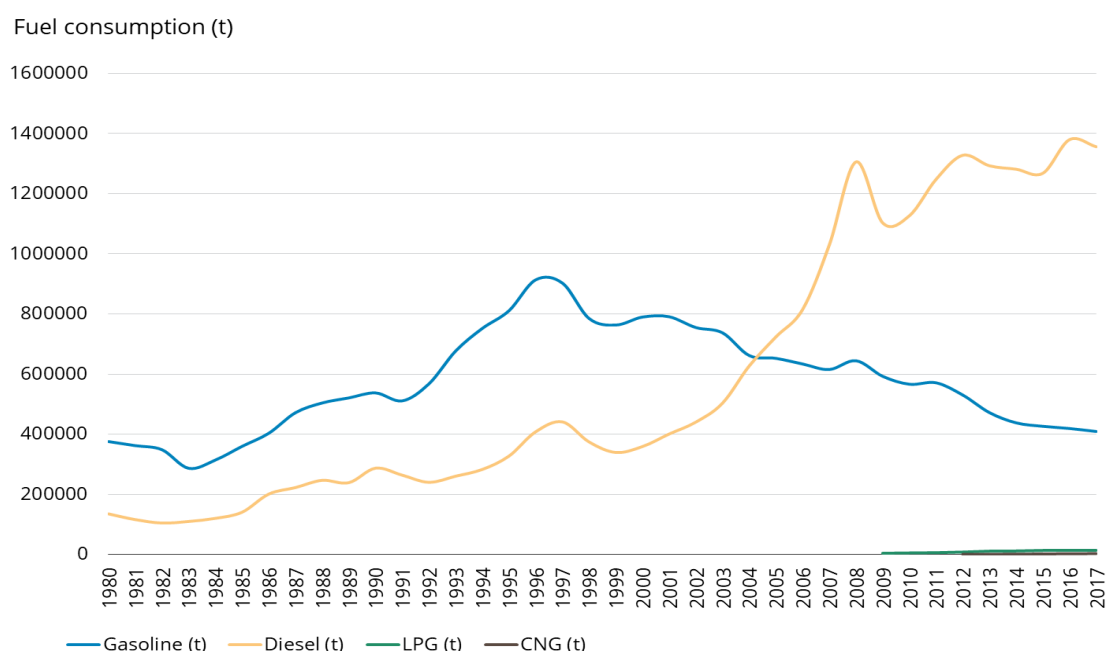
Three driving modes are individualized in accordance with COPERT 4 methodology: urban, rural and highway. For each specific driving mode average speeds has to be set by vehicles type whereas vehicle exhaust emissions and fuel consumption are strongly dependent on speed. Speeds in specific driving modes have been assessed on the basis of the Road Transport: Speed Data of the Republic of Slovenia publication, published by the Ministry of Transport. The values used are shown in the Annex 1 to the IIR (Table 1.4: Road transport: Speed data 1980–2017).

### Fuel Consumption

Statistical data on the total volume of fuel consumed in the Republic of Slovenia is obtained

from the SORS. From the total volume of fuel sold, the consumption in the fields of agriculture, forestry and construction has been excluded. Diesel, gasoline, liquefied petroleum gas (LPG) and compressed natural gas (CNG) have been used as fuels in road transportation.

As shown in Figure 3.3.1.2 the total fuel consumption in road transport began to grow markedly in the following two periods: during the years 1991-1997 due to fuel being sold to foreigners as a consequence of lower fuel prices in Slovenia, and during the years 1999-2008. During the years 2000-2008 an extensive growth in usage of diesel fuel can be observed. 2005 sale of diesel fuel exceeded the sale of gasoline. In 2009 a significant decline of gasoline and diesel consumption was observed. In comparison with the year 2008 consumption of gasoline dropped for 8 % and diesel for 16 %. Lower consumption of fuel was due to the world economic crisis. In the years 2011 and 2012 fuel consumption was on the rise again and slowly approaching pre-crisis values, but in the period 2013-2017 slightly lower fuel consumption could be observed. In 2017, the fuel use shares for diesel and gasoline were about 76 % and 23 %, respectively. The share of LPG was below 0,8 %. CNG was reported for the first time in 2012. It is mostly used in buses. Share of CNG is only 0,1 %.



**Figure 3.3.1.2 Fuel consumption in road transport for 1980–2017**

As shown in Figure 3.3.1.3 and Figure 3.3.1.4, passenger cars represent the most fuel-consuming vehicle category, followed by heavy duty trucks, light duty vehicles, buses, motorcycles and mopeds, in decreasing order. Fuel consumption for gasoline passenger cars dominates the overall gasoline consumption trend. The development in diesel fuel consumption in recent years is characterised by increasing fuel use for diesel passenger cars and heavy duty trucks, while the fuel use for buses and light duty vehicles is less distinctive. Due to transparency fuel consumption by types of vehicles is shown in the table in the Annex 1 to the IIR (Table 1.5: Road transport: Fuel Consumption by types of vehicle 1980 – 2017).

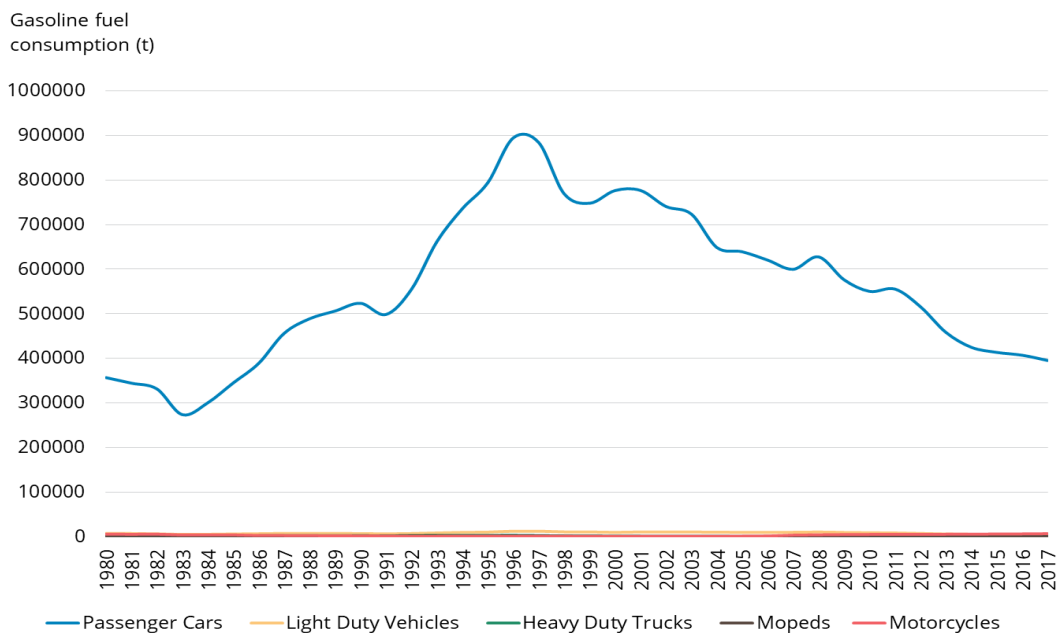


Figure 3.3.1.3 Gasoline fuel consumption per vehicle type for road transport 1980–2017

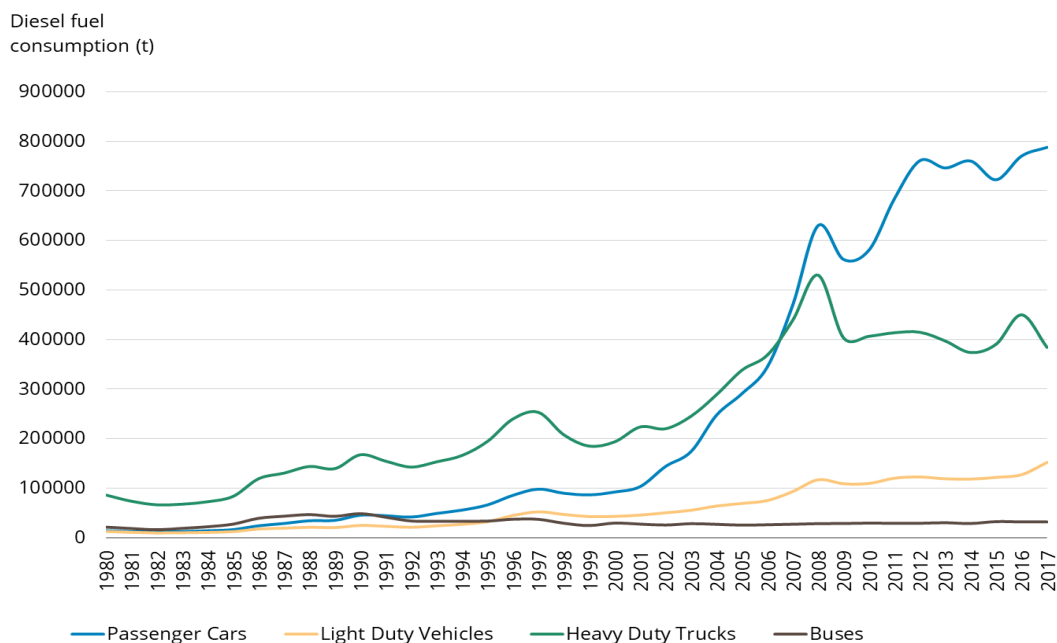


Figure 3.3.1.4 Diesel fuel consumption per vehicle type for road transport 1980–2017

In 2017, fuel consumption shares for diesel passenger cars, diesel heavy duty trucks and gasoline passenger cars were about 44, 22, and 22 %, respectively (Figure 3.3.1.5).

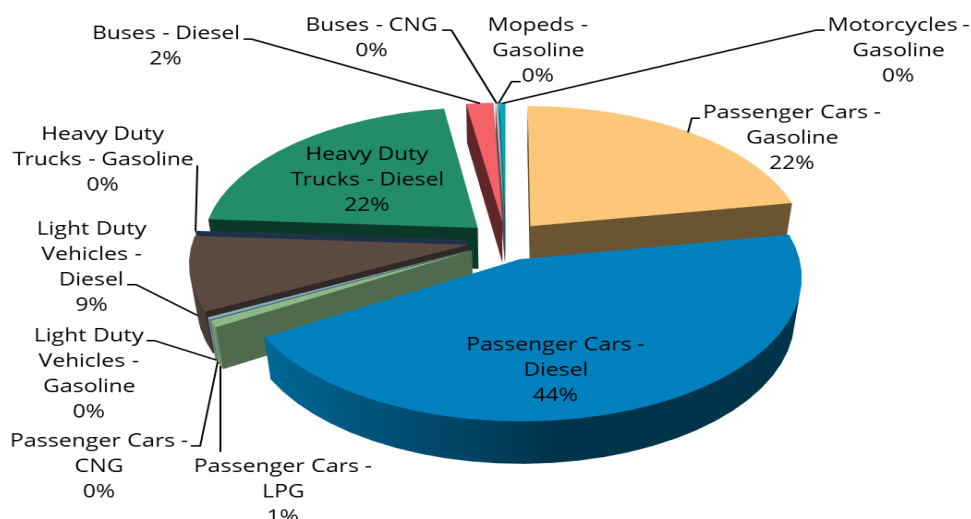


Figure 3.3.1.5 Fuel consumption share per vehicle type for road transport in 2017

### Fuel Characteristics

Sulphur and lead content of liquid fuels and monthly values of fuel volatility (RVP – Reid Vapour Pressure) were taken from Slovene national legislation relating quality of liquid fuels. Leaded gasoline was removed from the market in 2002. All the other physical and chemical data used was proposed as default values by the COPERT 4.

RVP values used were 70 kPa for winter period (1 October – 30 April) and 60 kPa for summer period (1 May – 30 September). The sulphur and lead contents were set as presented in Table 3.3.1.2 and Table 3.3.1.3.

Table 3.3.1.2 Levels of sulphur content in gasoline and diesel fuel

Fuel	Period	Sulphur [% wt]
Gasoline Leaded	1980-1994	0,1
	1995-2001	0,05
Gasoline Unleaded	1986-1994	0,1
	1995-2001	0,05
	2002-2004	0,015
	2005-2008	0,005
	2009-2017	0,001
Diesel	1980-1994	1
	1995	0,25
	1996-2001	0,20
	2002-2004	0,035
	2005-2008	0,005
	2009-2017	0,001

**Table 3.3.1.3 Levels of lead content in gasoline**

Fuel	Period	Lead [g/l]
Gasoline Leaded	1980-1994	0,6
	1995	0,4
	1996-2001	0,15
Gasoline Unleaded	1986-1994	0,026
	1995-2001	0,013
	2002-2017	0,005

### Monthly minimum and maximum air temperatures

Meteorological data necessary for evaporative emission calculation (annual average minimum temperature and maximum temperature) was obtained from Slovenian Environment Agency. Data for Ljubljana was taken into consideration with the assumption that it is representative enough for the whole Slovenia. Data are publicly available on Slovenian Environment Agency's website.

### Other input data

The average trip length (Ltrip) value corresponds to the mean distance covered in trips started with an engine of ambient temperature (cold start). Mean daily trip distance was set at 12 km in accordance with the recommendation of the COPERT 4. Ltrip value is introduced for the calculation of the Beta value which represents the fraction of the monthly mileage driven before the engine and any exhaust components have reached their nominal operation temperature. Beta values calculated according to the COPERT 4 methodology were used.

All the other required input data used for calculation of emissions using COPERT 4 program were default COPERT 4 data as well.

### Emission factors

All emission factors for NO<sub>x</sub>, SO<sub>x</sub>, NMVOC, NH<sub>3</sub>, PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC, CO, Pb, Cd, Cr, Cu, Ni, Se, Zn, dioxins/furans and PAHs (benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene, indeno(1,2,3-cd)pyrene), HCB, PCB used in the emission inventory for the whole period 1980 - 2017 are default emission factors offered by the COPERT 4 (version 11.4).

Emission factors for particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC) from automobile tyre and brake wear and road abrasion have been obtained from EMEP/EEA air pollutant emission inventory guidebook, 2016, Chapters: 1.A.3.b.vi and 1.A.3.b.vii. Road transport: automobile tyre and brake wear, automobile road abrasion, page 13-14, Table 3-1 and Table 3-2 for the whole period 2000 - 2017.

Emissions factors for calculation emissions of heavy metals (Pb, Cd, Cu, Cr, Ni Se, Zn) from automobile tyre and brake wear period for 1990 - 2017 are default emission factors offered by the COPERT 4 (version 11.4).

Emission factors for calculating exhaust particulate emissions represent together filterable and



condensable emissions. Exhaust emissions are considered to be PM<sub>2.5</sub>

There is no information for automobile tyre and brake wear and road abrasion whether emission factors represent filterable TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emissions or total (filterable and condensable) emissions.

### Emissions of SO<sub>x</sub>, NO<sub>x</sub>, CO, NMVOC, NH<sub>3</sub> and PM

From 1980 to 2017 the road transport emissions of SO<sub>x</sub> and CO have decreased by 99 and 91 %. In the same period, the emissions of NO<sub>x</sub> have increased by 6 %. Emissions of NMVOC have decreased by 90 % from 1990 to 2017 and emissions of NH<sub>3</sub> have increased by 2003 % from 1986 to 2017. From 2000 to 2017 emissions of exhaust PM have decreased by 23 %, while emissions of BC have increased by 2 %. Due to the world economic crises and consecutively smaller fuel consumption emissions of all pollutants considerably decreased in 2009. Decreasing trend is observed for the period 2010 - 2015 as well due to smaller fuel consumption and improved vehicle technologies. In 2016 the change of trend is observed. Sale of fuel was on the rise again. In 2017 a drop of emissions occurred due to smaller fuel consumption. Sale of gasoline and diesel decreased for about 2 %. Lower emissions compared to previous years were also due to bigger share of Euro 6 passenger cars and light duty vehicles and heavy duty trucks.

The gradual lowering of the sulphur content in diesel and gasoline fuel has given rise to a substantial decrease in the road transport emissions of SO<sub>x</sub>. In 1995, the sulphur content was reduced from 0,1 % (wt) to 0,05 % (wt) for gasoline and from 1 % (wt) to 0,25 % (wt) for diesel. The next clearly indicated emission drop occurred in 2002, when another substantial reduction in sulphur content in gasoline and diesel fuel were carried out. The last reduction of sulphur content in gasoline and diesel was performed in 2009. Sulphur content was reduced to 0,001 % (wt) in both fuels (Figure 3.3.1.6)

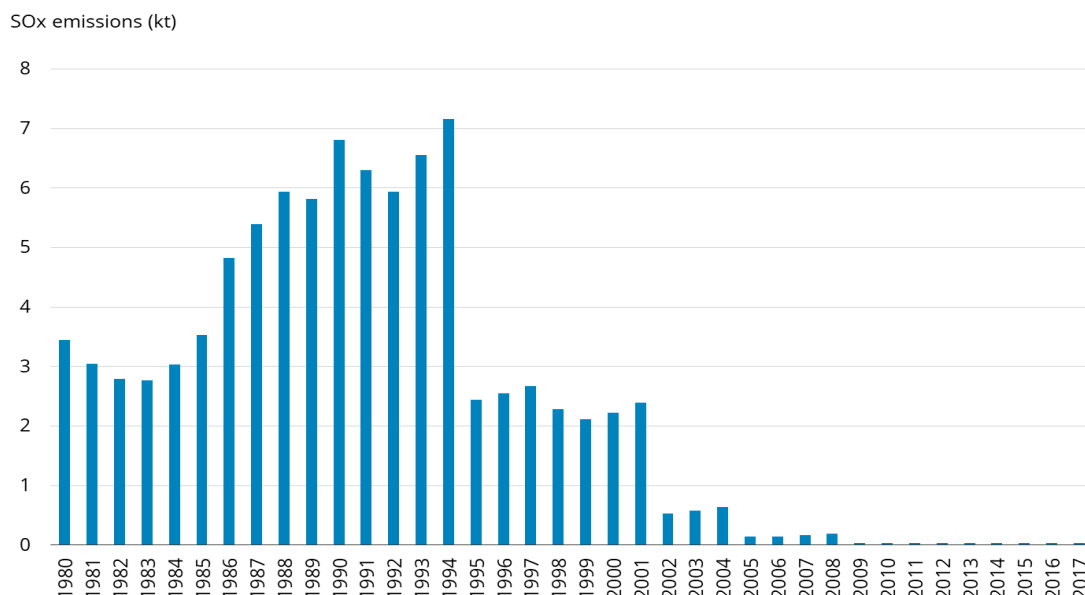
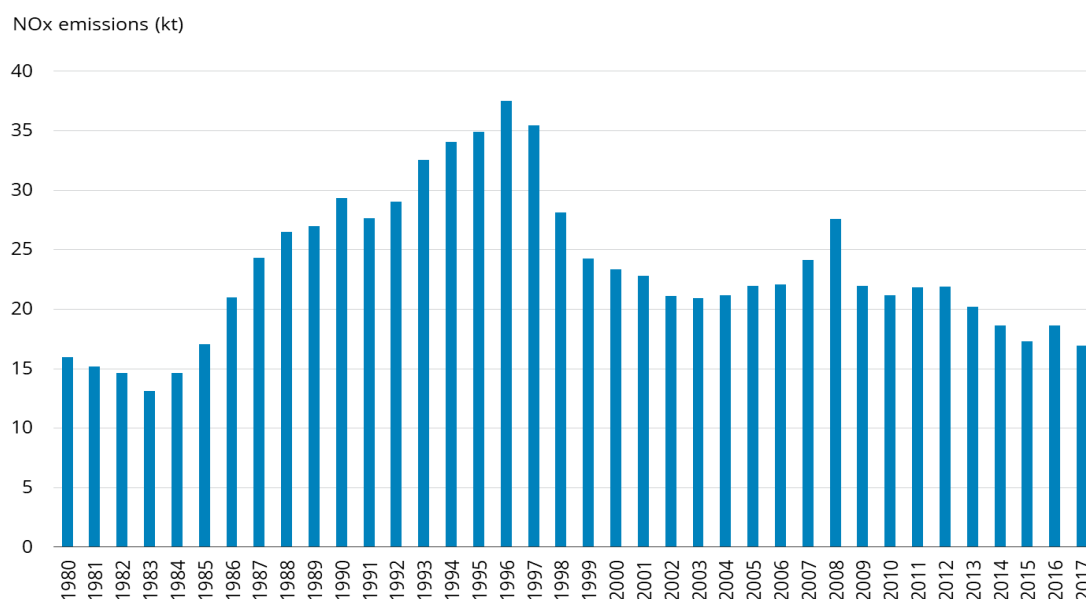


Figure 3.3.1.6 SO<sub>x</sub> emissions (kt) in road transport 1980–2017

NO<sub>x</sub> emissions have shown a steady decreasing tendency since the introduction of emission efficiently Euro 2 and Euro 3 catalyst cars into the Slovene fleet (introduced in 1997 and 2001, respectively). The positive effect of implementation of the stricter EURO standards has been made to no avail, due to the increased motor fuel consumption. Lower emissions in 2013, 2014 and 2015 are due to lower fuel consumption and introduction of EURO VI heavy duty trucks and Euro 6 passenger cars in national fleet. Increase in 2016 emissions was due to bigger diesel consumption compared to previous years. Lower emissions in 2017 were due to bigger share of Euro 6 passenger cars, light duty vehicles and heavy duty trucks. Small fuel consumption in 2017 is also a reason for drop of emissions (Figure 3.3.1.7).



**Figure 3.3.1.7 NO<sub>x</sub> emissions (kt) in road transport 1980–2017**

NM VOC and CO emissions have decreased in the last few years due to the growing share of vehicles that meet the stricter EURO standards. NM VOC and CO emission drops are also due to the decreasing share of gasoline passenger cars, as well as the decline in gasoline evaporation (Figure 3.3.1.8 and Figure 3.3.1.9).

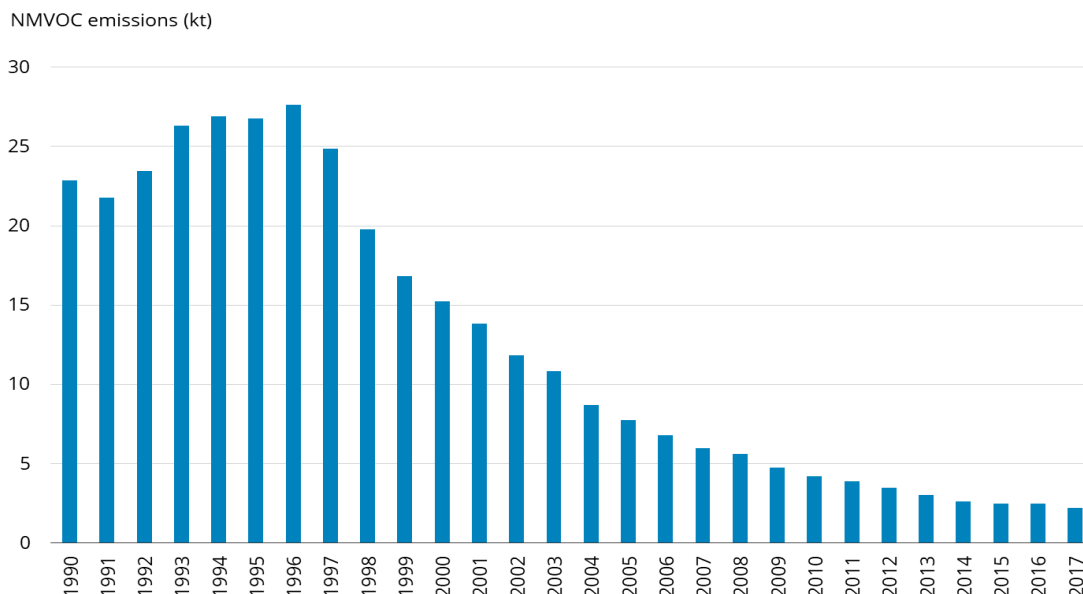


Figure 3.3.1.8 NMVOC emissions (kt) in road transport 1990–2017

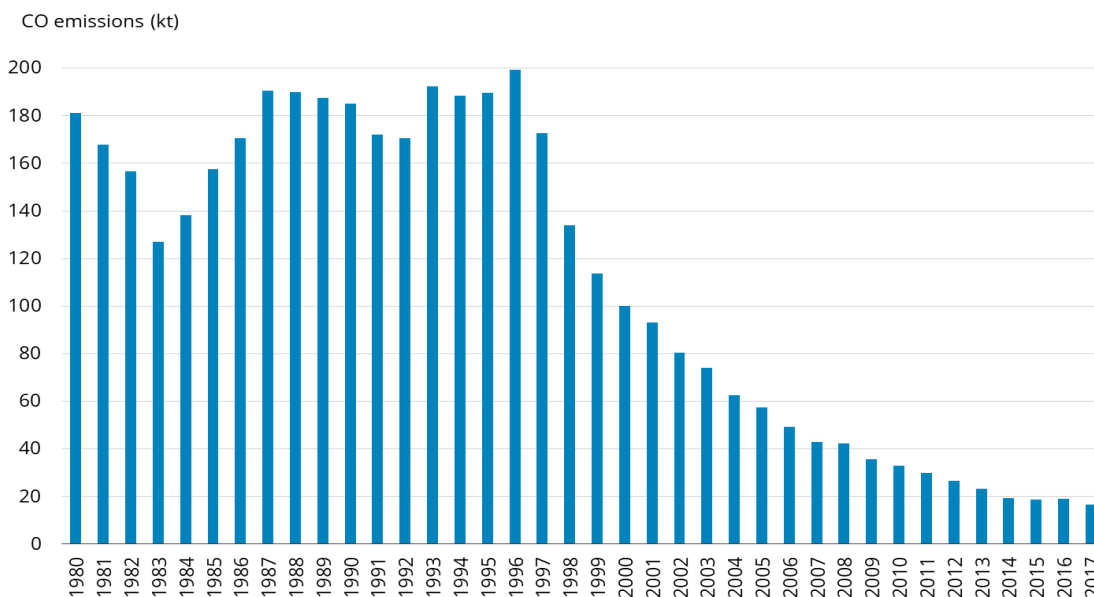


Figure 3.3.1.9 CO emissions (kt) in road transport 1980–2017

NH<sub>3</sub> emissions have increased rapidly from the year 1993 onward. The significant emission growth is related to the growth in the number of gasoline passenger cars fitted with catalyts. These produce ammonia as a by-product of the catalytic process that reduces emissions of nitrogen oxides. In the last few years the growth in emissions has stabilised, mostly due to the growth in the share of diesel passenger cars and consequently due to greater diesel fuel consumption (Figure 3.3.1.10).

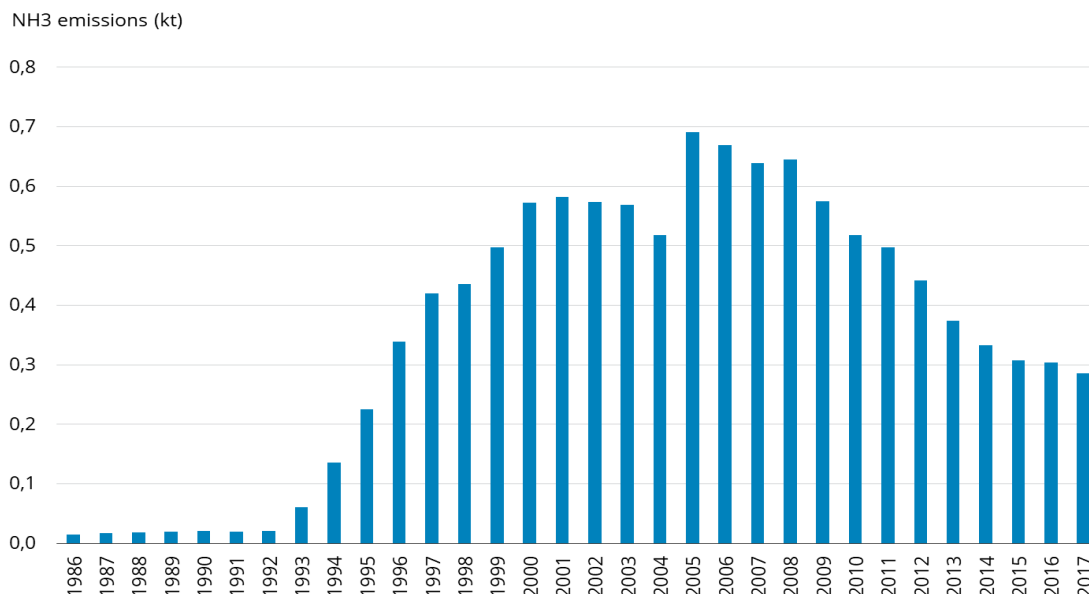


Figure 3.3.1.10 NH<sub>3</sub> emissions (kt) in road transport 1986–2017

Particulate emissions in the vehicle exhaust mainly fall in the PM<sub>2.5</sub> size range. Therefore, all PM emission corresponds to PM<sub>2.5</sub>. PM emission reduction has been achieved due to the growing share of vehicles that meet the stricter EURO standards. Also fuel refinements (mainly sulphur content reduction) played an important role in PM emission (Figure 3.3.1.11).

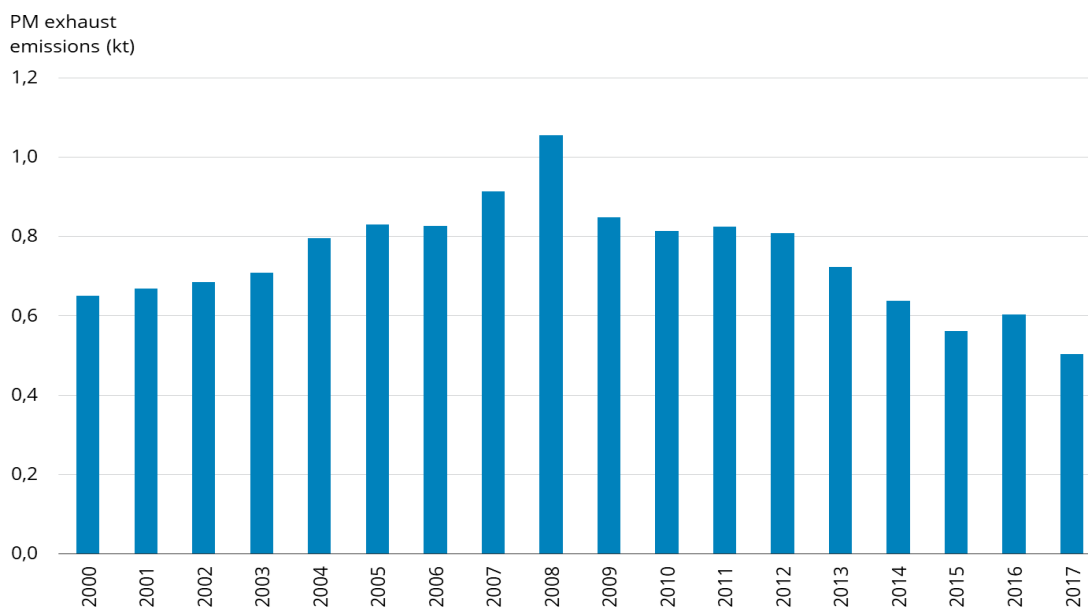
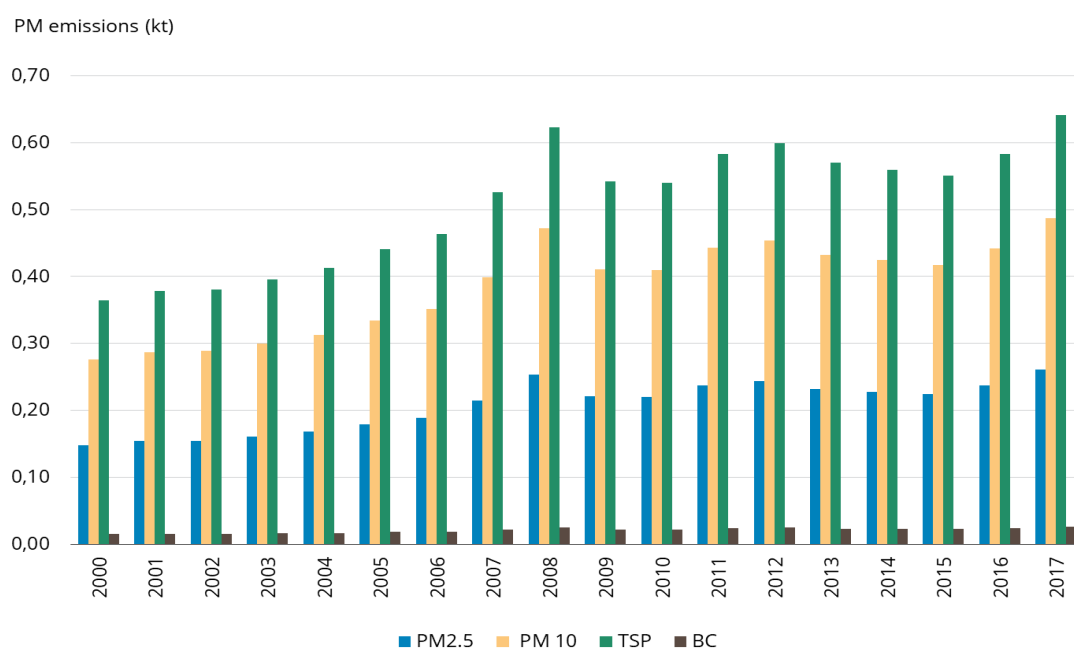
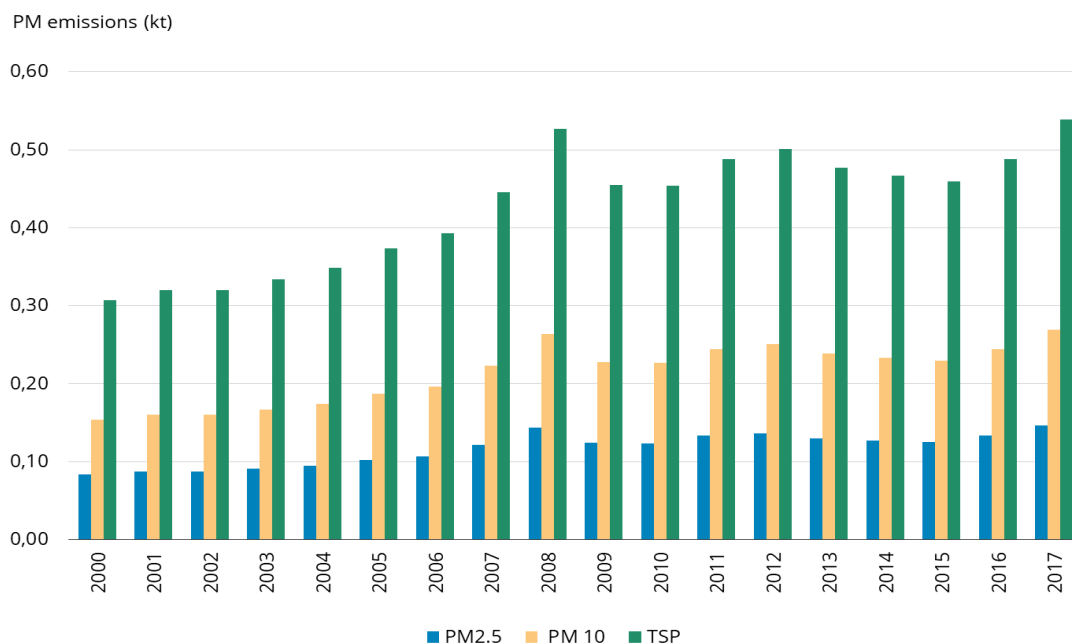


Figure 3.3.1.11 Exhaust PM emissions (kt) in road transport 2000–2017

Airborne particles are produced as a result of the interaction between a vehicle's tyres and the road surface, and also when the brakes are applied to decelerate the vehicle. Those particles emitted directly as a result of the wear of surfaces and not those resulting from the resuspension of previously deposited material. A jump of particulates emission from road vehicle tyre, brake wear and road abrasion in the year 2008 was a consequence of bigger fuel consumption and vehicle kilometres driven. In 2009 a significant decline of gasoline and diesel consumption was observed. In comparison with the year 2008 consumption of gasoline dropped for 8 % and diesel for 16%. This was reflected in decline of PM emissions. Lower consumption of fuel was due to the world economic crisis. Emissions for particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC) from automobile tyre and brake wear and road abrasion depend on total mileage driven and vehicle category (Figure 3.3.1.12, Figure 3.3.1.13 and Figure 3.3.1.14).

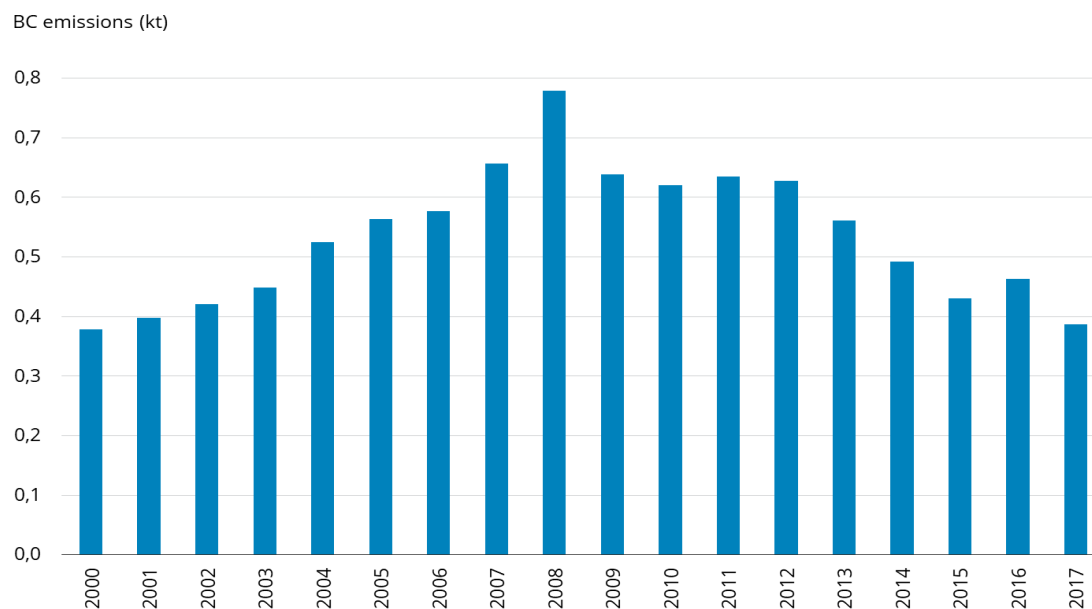


**Figure 3.3.1.12 PM emissions from road vehicle tyre and brake wear (kt) in road transport 2000–2017**



**Figure 3.3.1.13 PM emissions from road surface wear (kt) in road transport 2000–2017**

Emissions of black carbon (BC) mostly origin from vehicle exhaust, but smaller part also from automobile tyre and brake wear. Emissions of BC follow PM<sub>2.5</sub> emissions (Figure 3.3.1.14).



**Figure 3.3.1.14 BC emissions from road transport 2000–2017**

## **Emissions of Pb, Cd, PAHs, HCB, PCB, Dioxins and Furans**

From 1990 to 2017 the road transport emissions of Pb and PCB have decreased by 99 and 53 %. In the same period, the emissions of Cd, HCB, dioxins/furans and PAHs have increased by 103, 131, 71, 255 %, respectively. Road transport emissions of Pb, Cd, PAHs, dioxins/furans, HCB, PCB for the period 1990-2017 are shown in Figure 3.3.1.16 - Figure 3.3.1.21. Emissions of heavy metals presented in Figures comprise exhaust and non-exhaust emissions.

Pb emissions have decreased greatly from 1995-2017. The lowering is due to stricter legislation relating the content of Pb in gasoline fuel. Emissions of Cd have increased in the last few years due to bigger fuel consumption. Total emissions of four PAHs (indeno(1,2,3-cd)pyrene, benzo(k)fluoranthene, benzo(b)fluoranthene, benzo(a)pyrene) have been increasing due to changes in fleet vehicles. Total emissions of dioxins and furans have been decreasing due to growth in the share of diesel passenger cars. Increase of emissions in 2008 was due to bigger fuel consumption. Due to the world economic crises and consecutively smaller fuel consumption emissions of all pollutants considerably decreased in 2009. Decreasing trend is observed for the period 2010-2015 as well due to smaller fuel consumption and improved vehicle technologies. In 2016 the change of trend is observed. Sale of fuel was on the rise again. Lower emissions in 2017 were due to bigger share of Euro 6 passenger cars, light duty vehicles and heavy duty trucks. Small fuel consumption in 2017 is also a reason for drop of emissions.

According to 2018 in-depth EU NECD review emissions of Pb and Cd from automobile tyre and brake wear have been estimated for the whole period 1990-2017. Copert 4 (version 11.4) have been used for estimation of non-exhaust emissions.

Emissions of additional heavy metals (Cr, Cu, Ni, Se and Zn) have been calculated and included into the national inventory for the first time. Emissions of Cr, Cu, Ni, Se and Zn were introduced for the whole period 1990-2017. Copert 4 (version 11.4) have been used for estimation of exhaust emissions as well as non-exhaust emissions. The same methodology was used for emissions estimation for priority and additional heavy metals.

Emissions of Hg and As have not been calculated since Copert model does not provide emission factors for those two heavy metals.

Emissions of Cr, Cu, Ni, Se and Zn have been increased between 1990 and 2017 by 145, 139, 119, 141 and 114 %, respectively. A jump of heavy metals emissions in the year 2008 was due to bigger fuel consumption. In 2009 a significant decline of gasoline and diesel consumption was observed. In comparison with the year 2008 consumption of gasoline dropped for 8 % and diesel for 16%. This was reflected in decline of emissions. Road transport emissions of Cr, Cu, Ni, Se and Zn for the period 1990 – 2017 are shown in Figure 3.3.1.22 - Figure 3.3.1.26.

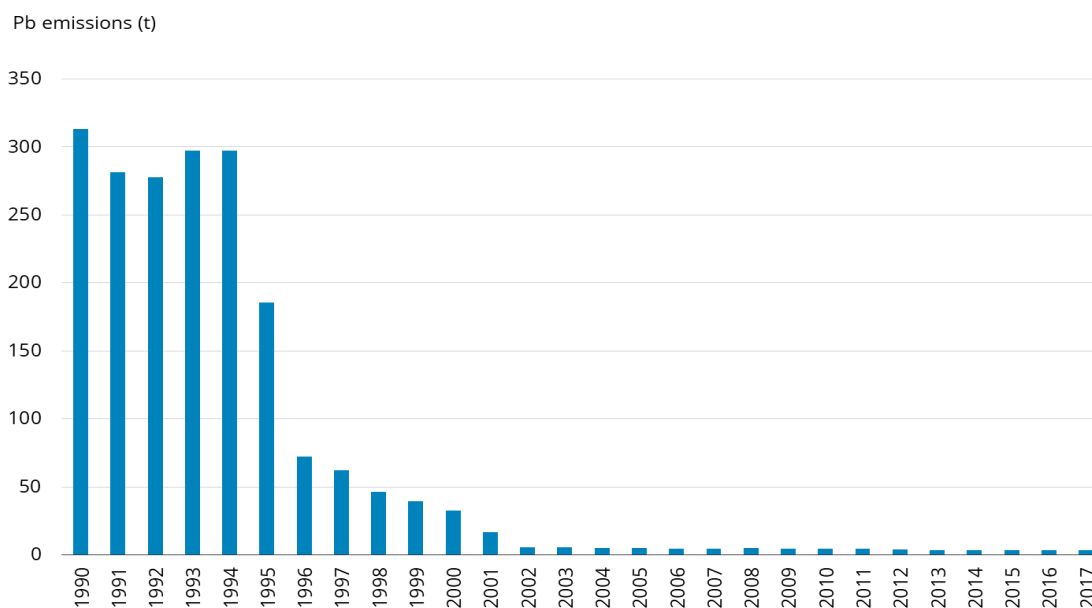


Figure 3.3.1.16 Pb emissions (t) in road transport 1990–2017

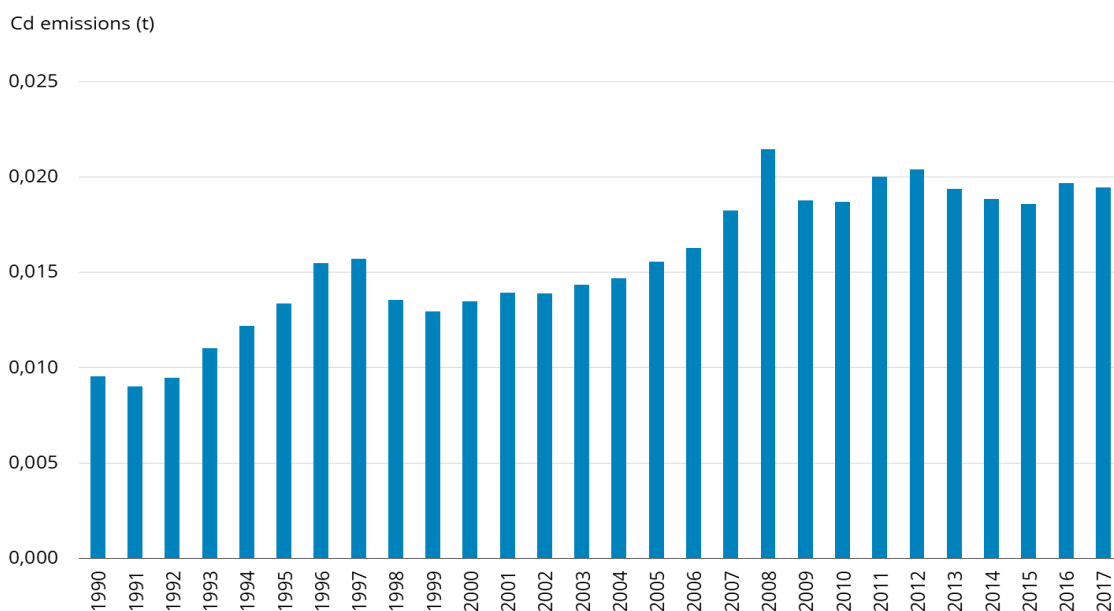


Figure 3.3.1.17 Cd emissions (t) in road transport 1990–2017



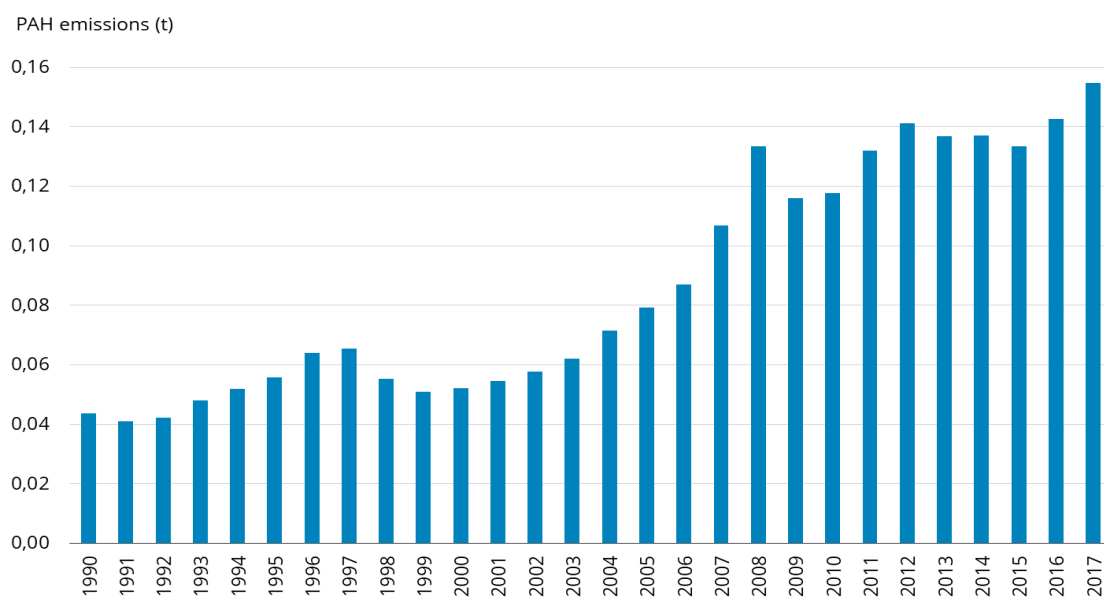


Figure 3.3.1.18 PAHs emissions (t) in road transport 1990–2017

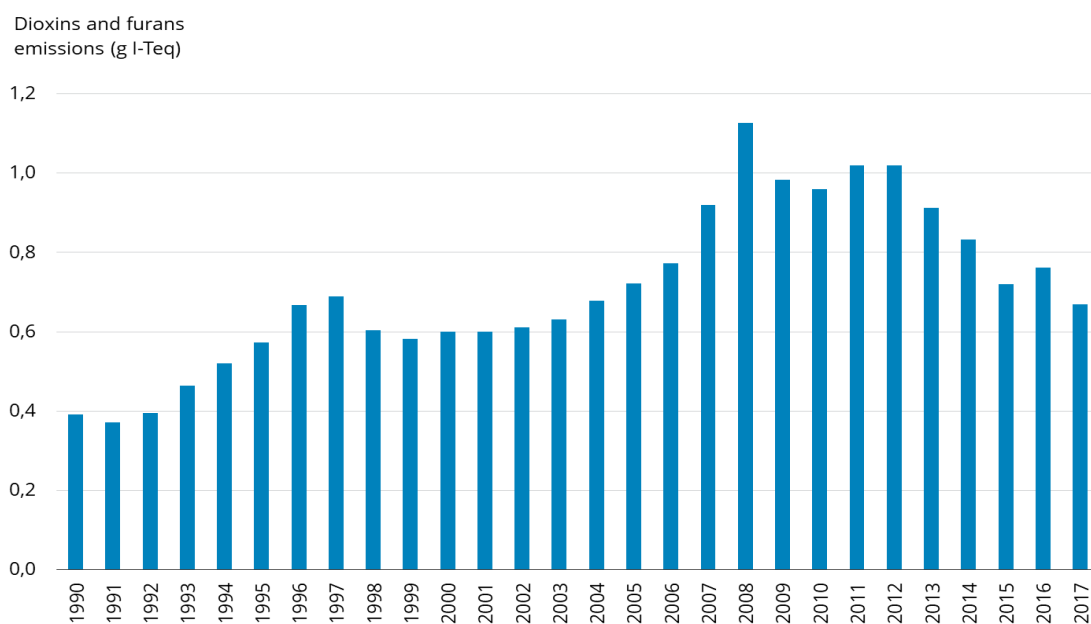


Figure 3.3.1.19 Dioxins/Furans emissions (g I-Teq) in road transport 1990–2017

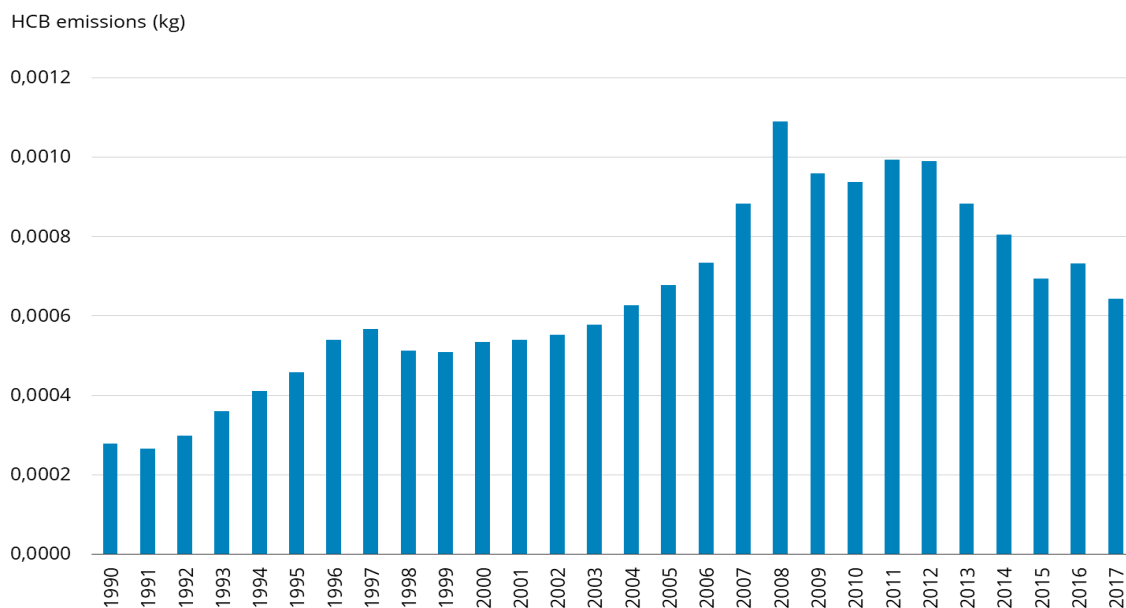


Figure 3.3.1.20 HCB (kg) in road transport 1990–2017

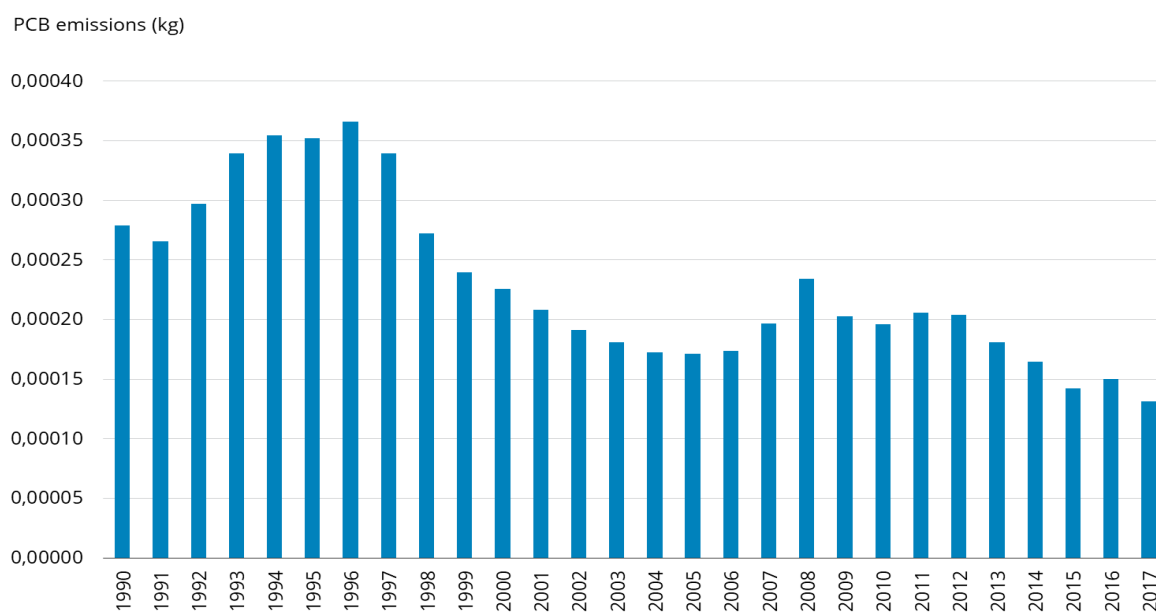


Figure 3.3.1.21 PCB (kg) in road transport 1990–2017

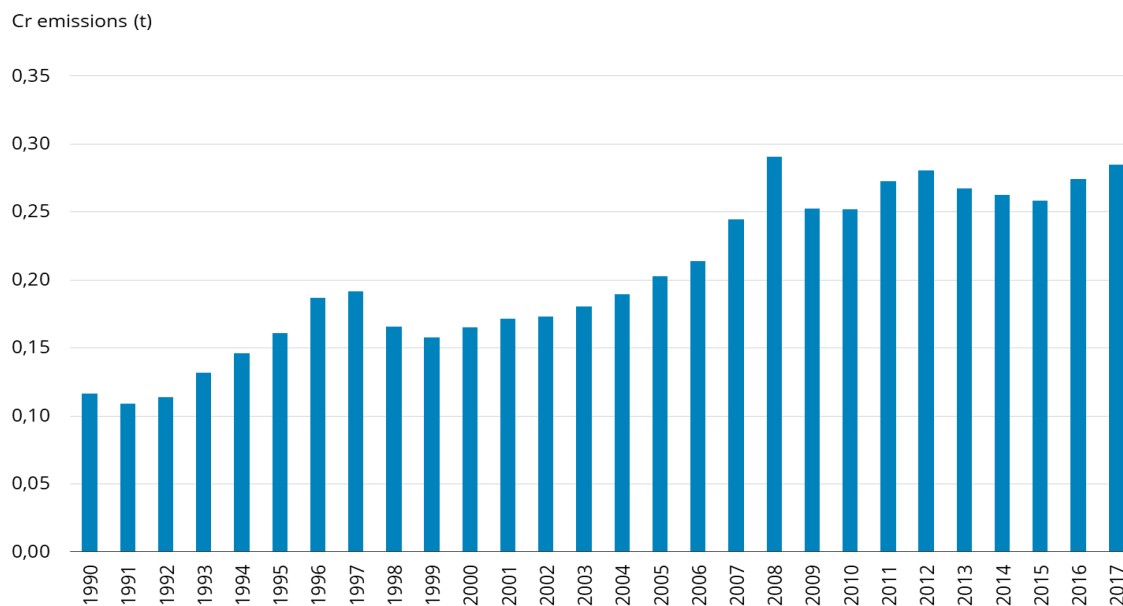


Figure 3.3.1.22 Cr (t) in road transport 1990–2017

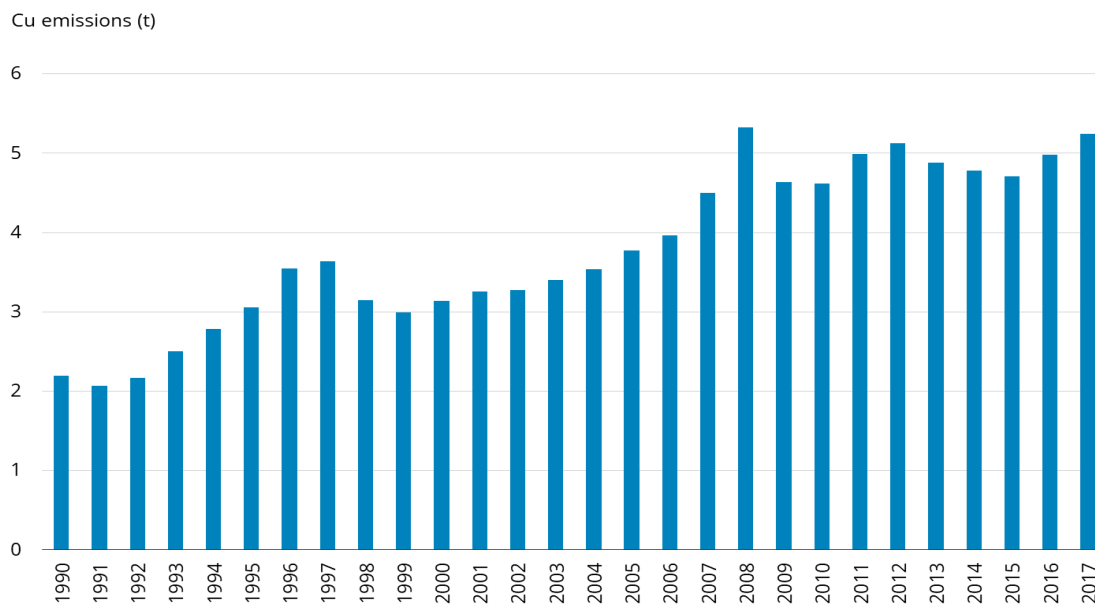


Figure 3.3.1.23 Cu (t) in road transport 1990–2017

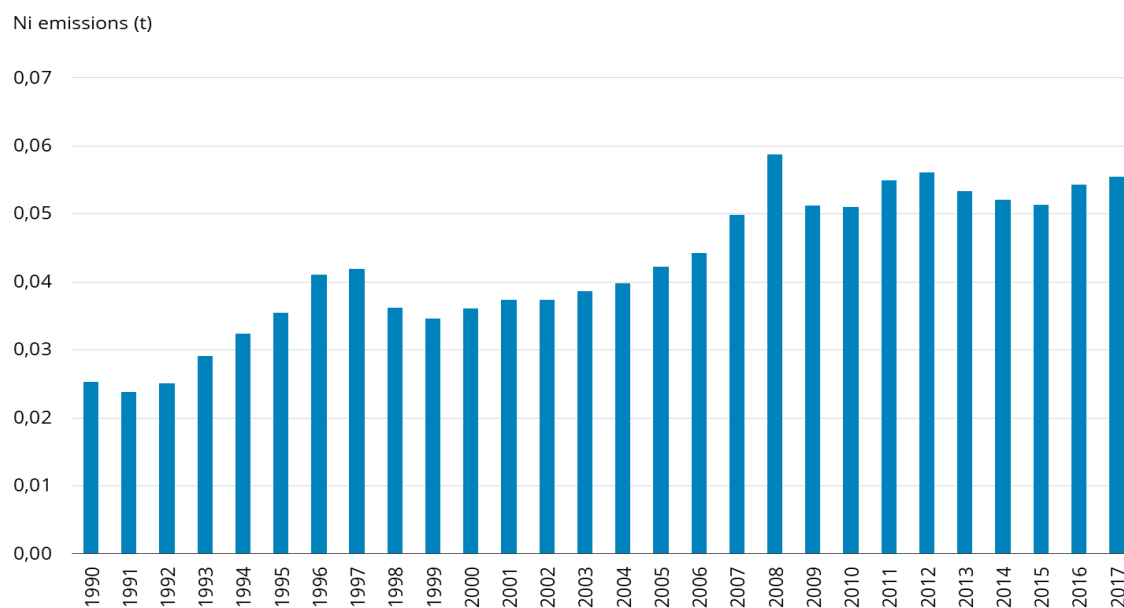


Figure 3.3.1.24 Ni (t) in road transport 1990–2017

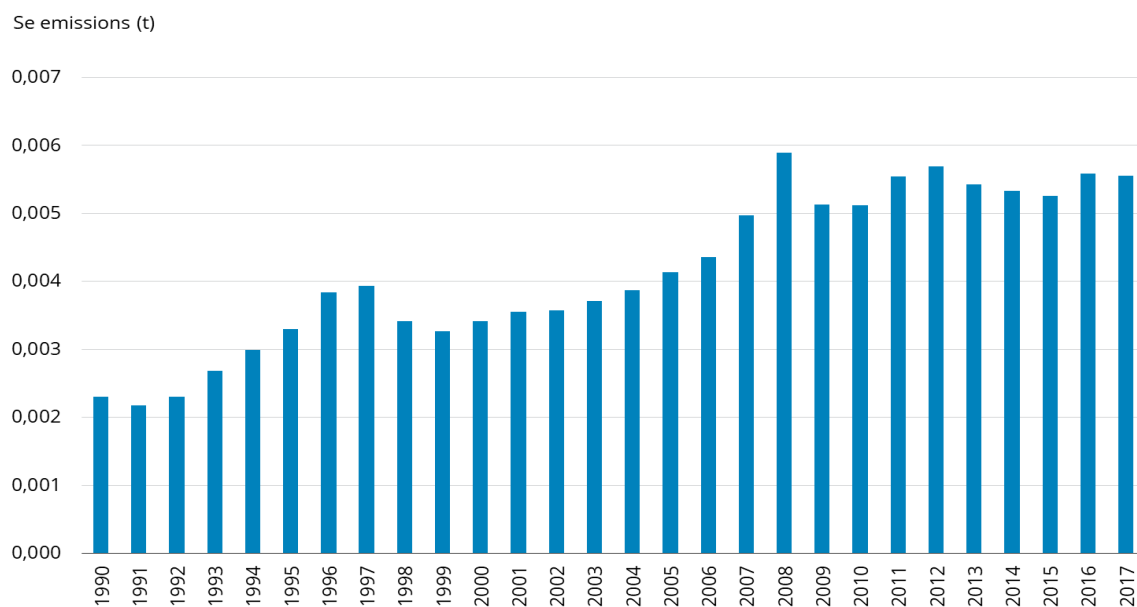
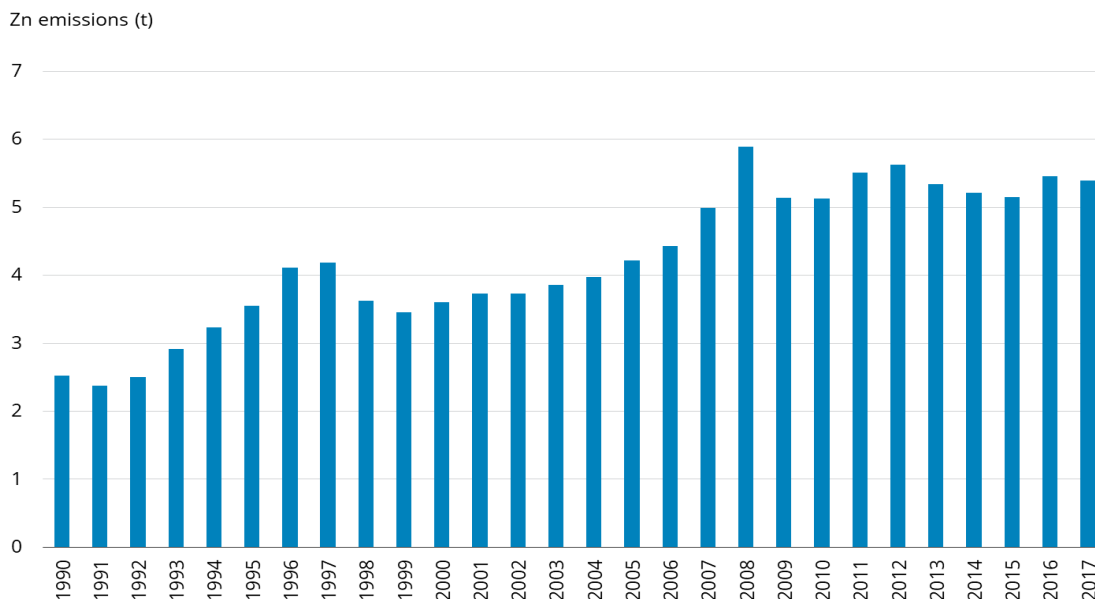


Figure 3.3.1.24 Se (t) in road transport 1990–2017



**Figure 3.3.1.24 Zn (t) in road transport 1990–2017**

### Recalculations

According to 2018 in-depth EU NECD review emissions of Pb and Cd from automobile tyre and brake wear have been estimated for the whole period 1990-2017. Copert 4 (version 11.4) have been used for estimation of non-exhaust emissions.

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017. Exhaust and non-exhaust emissions have been estimated.

Emission of SO<sub>x</sub> was recalculated for 2016 due to error in filling the NFR reporting table.

### Category-Specific QA/QC and Verification

Examination of input data, the model calculation and the data reported in NFR tables as part of QC/QC procedure was performed. Error in SO<sub>x</sub> emissions for 2016 was found. Mistake was corrected. External experts are checked the data.

### Planned improvements

We are planning to use new Copert 5 model for emission calculation from road transport in next two years.

### 3.3.2 Railways

NFR Code 1A3c

#### Introduction

Exhaust emissions from railways arise from the combustion of liquid fuels in diesel engines, and solid or liquid fuels in steam engines to provide propulsion. The principal pollutants are those from diesel engines, similar to those used in road transport. In the year 2017 railways mostly contributed to the total NO<sub>x</sub> (1,4 %) and to a lesser extent to other pollutants.

#### Methodology

To estimate emissions from the railways the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (kg/t) or (g/GJ)

In case of EF expressed in the unit g/GJ net calorific value (NCV) of fuel is needed for emission calculation.

#### Activity data

The main source of emissions is a consumption of diesel. The consumption of coal in railway transportation was small, from 0 to 646 t. This coal was used in only one "archaic" steam driven locomotive which is almost 100 years old. The specified data have been obtained from Statistical Office of the Republic of Slovenia (SORS).

There were no data available on consumption of diesel and brown coal used in railway sector before 1986. Activity data for the period 1980-1985 have been estimated.

Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.6: Fuel Consumption: Railways).

#### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used for emissions calculation.

**Table 3.3.2.1 Emission factors for diesel used for emission calculation and references**

Pollutant	Diesel	Unit	References
NO <sub>x</sub>	52,4	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
SO <sub>x</sub>	Values used for road transport (Table 3.3.1.2)	kg/t	Slovene national legislation relating quality of liquid fuels
CO	10,7	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1

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<b>NM VOC</b>	4,65	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>NH<sub>3</sub></b>	0,007	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>PM<sub>2.5</sub></b>	1,37	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>PM<sub>10</sub></b>	1,44	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>TSP</b>	1,52	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>BC</b>	0,8905	kg/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Cd</b>	0,01	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Cr</b>	0,05	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Cu</b>	1,7	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Ni</b>	0,07	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Se</b>	0,01	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Zn</b>	1	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Benzo(a)pyrene</b>	0,03	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Benzo(b)fluoranthene</b>	0,05	g/t	Emission Inventory Guidebook, 2016, 1.A.3.c Railways, pg 8, Table 3-1
<b>Benzo(k)fluoranthene</b>	0,0344	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
<b>Indeno(1,2,3-cd)pyrene</b>	0,0079	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
<b>Pb</b>	0,052	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-10

Table 3.3.2.2 Emission factors for sub-bituminous coal used for emission calculation and references

Pollutant	Coal	Unit	References
<b>NO<sub>x</sub></b>	209	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>SO<sub>x</sub></b>	820	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>CO</b>	8,7	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>NM VOC</b>	1,0	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>PM<sub>2.5</sub></b>	3,4	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>PM<sub>10</sub></b>	7,7	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>TSP</b>	11,4	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>BC</b>	0,0748	g/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Cd</b>	0,9	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Hg</b>	1,4	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Pb</b>	7,3	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2

<b>As</b>	7,1	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Cr</b>	4,5	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Cu</b>	7,8	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Ni</b>	4,9	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Se</b>	23	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Zn</b>	19	mg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Dioxins/ Furans</b>	10	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>HCB</b>	6,7	microg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Benzo(a)pyrene</b>	0,7	microg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Benzo(b)fluoranthene</b>	37	microg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Benzo(k)fluoranthene</b>	29	microg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2
<b>Indeno(1,2,3-cd)pyrene</b>	1,1	microg/GJ	Emission Inventory Guidebook, 2016, 1.A.1 Energy industries, pg 16, Table 3-2

There is no information for diesel whether emission factors include or exclude condensable component. PM emission factors for coal used represent filterable PM emissions.

### Net calorific values

Data on NCV have been obtained from SORS.

**Table 3.3.2.3 NCV for brown coal and diesel used for emission calculation**

Fuel	NCV	Unit
Diesel	42,6	MJ/kg
Coal	11,64	MJ/kg

### Emissions

In the year 2017 railways mostly contributed to the national total NO<sub>x</sub> (1,4 %) and to a lesser extent to other pollutants. There is a strong increase in diesel consumption in 2014. The reason for this increase is a severe ice storm which destroyed electrical infrastructure for the supply of trains on the route Ljubljana - Koper in the February 2014. The repair was going on until the summer 2015. In meantime, the trains on this line were using diesel locomotives what resulted in the higher consumption of diesel oil in 2014 and relatively high consumption in 2015.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 were used for emissions calculation. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb and NMVOC were recalculated for the period 1990-2016. Recalculation of PM<sub>2,5</sub>, PM<sub>10</sub>, TSP, BC were performed for the period 2000-2016 and recalculation of SO<sub>x</sub> and NO<sub>x</sub> for the period 1980-2016.



## Future Improvements

No improvement is planned for next submission.

### 3.3.3 Aviation

Sectors covered in this chapter are:

NFR Codes:

1A3ai(i)	International aviation LTO (civil)
1A3aai(i)	Domestic aviation LTO (civil)
1A5b	Other, Mobile (including military, land based and recreational boats)

#### 3.3.3.1 International aviation LTO (civil)

NFR Code 1A3ai(i)

#### Introduction

In sector international aviation are included journeys where aircrafts depart from one country and arrive in another country. There is only one operative international airport in Slovenia (Aerodrom Ljubljana) where international airport traffic has been taking place. Exhaust emissions from international airport traffic aviation arise from the combustion of jet kerosene. The landing and take-off cycle includes all activities near the airport that take place below a height of 3 000 ft (914 m). This therefore includes taxi-in and -out, take-off, climb-out and approach-landing. Contribution to total national emissions for all pollutants is below 1 %.

#### Methodology

To estimate emissions from international aviation, the following methodology has been adopted:

$$E = m \times EF$$

E - emission (kg)

m - quantity of fuel combusted (t)

EF - emission factor per quantity of fuel (kg/t)

#### Activity data

Quantity of jet kerosene applied for emission calculation has been obtained from Statistical Office of the Republic of Slovenia (SORS). Amount of fuel used in 2017 was 23668 t. Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.7: Fuel Consumption: International aviation LTO (civil)).

#### Emission factors

Emission factors were calculated from annual fuel consumption obtained from Statistical Office of the Republic of Slovenia and emission factors for the landing and take-off cycle (LTO cycles)

as well as fuel consumption for certain aircraft type. LTO fuel consumption and emission factors for certain aircraft types were obtained from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23. Data for aircraft type A320 was used for emission calculation of individual gases.

**Table 3.3.3.1.1 Emission factors for jet kerosene used for emission calculation and references**

Pollutant	Jet kerosene	Unit	References
NO <sub>x</sub>	13,28	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
SO <sub>x</sub>	0,84	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
CO	10,1	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
NM VOC	1,81	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
PM <sub>2.5</sub>	0,08	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
PM <sub>10</sub>	0,08	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23

There is no information whether emission factors represent filterable PM<sub>10</sub> and PM<sub>2.5</sub> emissions or total (filterable and condensable) emissions.

### Recalculations

No recalculations have been performed since last submission.

### Future Improvements

No improvements are planned for next submission.

### 3.3.3.2 Domestic aviation LTO (civil)

NFR Code 1A3aii(i)

#### Introduction

Civil domestic aviation comprises journeys where aircrafts depart and arrive in the same country. In Slovenia there are a couple of small airports used for sport or tourist activities. Emissions are very low due to small amount of fuel used for these purposes. Contribution to total national emissions for all pollutants is below 1 %.

#### Methodology

To estimate emissions from civil aviation, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF - emission factor per quantity of fuel (kg/t)

#### Activity data

For domestic aviation gasoline and jet kerosene have been used. Quantity of fuel used has been obtained from SORS. Amount of aviation gasoline used in 2017 was 372 t. 183 t of jet kerosene was consumed as well. Fuel consumption for the whole period is shown in in the Annex 1 to the IIR (Table 1.8: Fuel Consumption: Domestic aviation LTO (civil)).

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 3.3.3.2.1 Emission factors for aviation gasoline used for emission calculation and references**

Pollutant	Fuel	Unit	References
NO <sub>x</sub>	4	kg/t	Emission Inventory Guidebook, 2016, Civil aviation (domestic, LTO), pg 21, Table 3-3
SO <sub>x</sub>	1	kg/t	Emission Inventory Guidebook, 2016, Civil aviation (domestic, LTO), pg 21, Table 3-3
CO	1200	kg/t	Emission Inventory Guidebook, 2016, Civil aviation (domestic, LTO), pg 21, Table 3-3
NM VOC	19	kg/t	Emission Inventory Guidebook, 2016, Civil aviation (domestic, LTO), pg 21, Table 3-3
Pb	0,033	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-10
Benzo(a)pyrene	0,0055	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-9
Benzo(b)fluoranthene	0,0079	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-9
Benzo(k)fluoranthene	0,0039	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
Indeno(1,2,3-cd)pyrene	0,0089	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8

**Table 3.3.3.2.2 Emission factors for jet kerosene used for emission calculation and references**

Pollutant	Jet kerosene	Unit	References
NO <sub>x</sub>	13,28	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
SO <sub>x</sub>	0,84	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
CO	10,1	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
NM VOC	1,81	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
PM <sub>2.5</sub>	0,08	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23
PM <sub>10</sub>	0,08	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-4, pg 23

There is no information whether emission factors represent filterable PM<sub>10</sub> and PM<sub>2.5</sub> emissions or total (filterable and condensable) emissions.

### Emissions

According to the Eurocontrol data, a small amount of jet kerosene has been used since 2005 in domestic aviation. Due to the increase in traffic in the summer time, some charter flights have been transferred to the Maribor airport. For this purpose, it was necessary to transfer the aircrafts from Ljubljana to Maribor and back. The amount of jet kerosene used for this purpose is very small. There are two peaks in the fuel consumption in the time series. One in 2005 is

connected to the inclusion of jet kerosene while we do not know the reason for the peak in 2011. However, the total amount of fuel is small and therefore even a small amount of fuel could influence a big relative change. High increase in fuel consumption in 2011 (+40%) was due to the increase of aviation gasoline for 87 tonnes and jet kerosene for 170 tonnes what are quite insignificant quantities.

### **Recalculations**

No recalculations have been performed since last submission.

### **Future Improvements**

No improvement is planned for next submission.

### **3.3.3.3 Other, Mobile (including military, land based and recreational boats)**

NFR Code 1A5b

#### **Introduction**

Military and police aircrafts and helicopters serve different purposes. Beside regular security operations and training activities they are also engaged in emergency medical service, intervention in natural disasters and mountain rescue operations. Emissions of main pollutants have been estimated from use of fuel in army and police air force fleet. Emissions do not contribute much (below 0,1 %) to the total emissions due to small amount of fuel used.

#### **Methodology**

To estimate emissions from army and police aviation the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (kg/t)

#### **Activity data**

Consumption of jet kerosene in Slovenian army and police for the period 1980 - 2017 has been obtained from both institutions. The consumption of fuel for helicopters and military flights was small due to small air force fleet. Consumption of jet kerosene in the year 2017 was 1318 t. Fuel consumption for the whole period is shown in in the Annex 1 to the IIR (Table 1.9: Fuel Consumption: Other, Mobile (including military, land based and recreational boats).

According to 2017 in-depth EU NECD review use of aviation gasoline was checked. All aviation gasoline sold in Slovenia is considered to be used for domestic aviation and the emissions are reported in category Domestic aviation, civil, LTO – (NFR 1.A.3.a.iii(i)). We have obtained this data only for last three years, however the data are not available for entire time series. According to data for 2015, 38,6 tonnes of aviation gasoline have been used in the army what is less than 10 per cent of total aviation gasoline used in this year. We believe that emissions from this

source are negligible and that disaggregation will not lead to a noticeable improvement.

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

**Table 3.3.3.3.1 Emission factors for jet kerosene used for emission calculation and references**

Pollutant	Jet kerosene	Unit	References
NO <sub>x</sub>	4,631	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
SO <sub>x</sub>	1,025	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
CO	33,9	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
NM VOC	2,331	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29

### Recalculations

No recalculations have been performed since last submission.

### Future Improvements

No improvement is planned for next submission.

## 3.3.4 Memo items - International bunker fuels

Sectors covered in this chapter are:

NFR Codes:

1A3di(i) International maritime navigation  
1A5c Multilateral operations

### 3.3.4.1 International maritime navigation

NFR Code 1A3di(i)

#### Introduction

Slovenia has only one international port "Luka Koper" but in the period 1980-2005 no ships had been refuelled in that port. Ships were mostly refuelled in the international waters by Italian ships under Panama flags. Since 2006 a small amount of heavy fuel oil has been reported as fuel sold to the international marine bunkers.

## Methodology

To estimate emissions from international maritime navigation the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (kg/t)

## Activity data

Quantity of heavy fuel oil used for emission calculation has been obtained from SORS for the period 2006-2017. Amount of bunker fuel oil used in 2017 was 157305 t. Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.10: Fuel Consumption: International maritime navigation/ International bunker fuels).

## Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

**Table 3.3.4.1.1 Emission factors for heavy fuel oil used for emission calculation and references**

Pollutant	Heavy fuel oil	Unit	References
NO <sub>x</sub>	79,3	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
SO <sub>x</sub>	1,0	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
CO	7,4	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
NM VOC	2,7	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
PM <sub>2.5</sub>	5,6	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
PM <sub>10</sub>	6,2	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
TSP	6,2	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
BC	0,672	kg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Cd	0,02	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Pb	0,18	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Hg	0,02	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
As	0,68	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Cr	0,72	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Cu	1,25	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Ni	32	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
Se	0,21	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13

<b>Zn</b>	1,2	g/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
<b>PCB</b>	0,57	mg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
<b>HCB</b>	0,14	mg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13
<b>Dioxins/ Furans</b>	0,0047	mg/t	Emission Inventory Guidebook, 2016, Navigation, Table 3-1, pg 13

There is no information whether emission factors represent filterable TSP, PM<sub>10</sub> and PM<sub>2.5</sub> emissions or total (filterable and condensable) emissions.

### Emissions

The emissions produced by navigation are a consequence of combusting the fuel in an internal combustion engine. According to revised guidelines for reporting emissions and projections data under the Convention (ECE/EB.AIR/122/Add.1, decisions 2013/3 and 2013/4) and EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 emissions resulting from international journeys are not included in national totals.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2006-2017.

### Future Improvements

No improvement is planned for next submission.

#### 3.3.4.2 Multilateral operations

NFR Code 1A5c

### Introduction

The Slovenian Armed Forces participate in multinational operations and missions in Afghanistan and Kosovo. Information on Slovenian cooperation in international operations is presented on web page:

<http://www.slovenskavojska.si/en/international-cooperation/international-operations-and-missions/>

### Methodology

To estimate emissions from international aviation (cruise) the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)  
 EF – emission factor per quantity of fuel (kg/t)

### Activity data

Quantity of jet kerosene used for emission calculation has been obtained from Slovenian army. According to the data from Slovenian Army about 15 % jet kerosene were used in international missions. Data are available for the period 1997-2017. Amount of jet kerosene used in multilateral operations in 2017 was 182 t. Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.11: Fuel Consumption: Multilateral operations/ International bunker fuels).

The amount of jet kerosene used in Slovene Army and Police is excluded from international aviation bunkers, and is reported under 1A5b Other, Mobile.

### Emission factors

**Table 3.3.4.2.1 Emission factors for jet kerosene used for emission calculation and references**

Pollutant	Jet kerosene	Unit	References
<b>NO<sub>x</sub></b>	4,631	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
<b>SO<sub>x</sub></b>	1,025	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
<b>CO</b>	33,9	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29
<b>NMVO</b>	2,331	kg/t	Emission Inventory Guidebook, 2016, Aviation, Table 3-11, pg 29

### Emissions

According to revised guidelines for reporting emissions and projections data under the Convention (ECE/EB.AIR/122/Add.1, decisions 2013/3 and 2013/4) and EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 emissions resulting from multilateral operations are not included in national totals.

### Recalculations

No recalculations have been performed since last submission.

### Future Improvements

No improvement is planned for next submission.



### **National navigation (Shipping); NFR Code 1A3dii**

Fuel used for small boats and yachts has been sold on four petrol stations at Adriatic coast (Izola Pier, Lucija Pier, Marina Koper and Piran Pier). These patrol stations are used for filling up road vehicles as well. A division between road and marine traffic is not possible. For this reason, we have reported all fuel in sub-sector road traffic. Notation Key "IE" (included elsewhere) was used for domestic water-borne navigation, since all fuel used for this sector was reported under 1A3b Road transport. We tried to obtain separate data, but we were not successful. Statistical Office, which is our main fuel data provider, does not provide those data. Since we have no possibilities for obtain reliable data, we are not planning to report data separately in next annual submission.

### **International inland waterways; NFR Code 1A3di(ii)**

Notation Key "NO" (not occurring) was used for this sector, since there is no emissions from international inland waterways in Slovenia.

### **International aviation cruise (civil): NFR Code 1A3ai(ii)-memo items**

Notation Key "IE" (included elsewhere) was used for International aviation cruise (civil), since all fuel used for this sector was reported under 1A3ai(i) International aviation LTO (civil). Overestimation of national emissions due to inclusion of memo category 1A3ai(ii) to national totals is below 1 % and does not have a significant impact on national inventory. Since this source is of less importance Tier 1 method was used for emission calculation. In addition, we have a national database (Emission inventory information system) which we use for calculation and reporting of greenhouse gas emissions and air pollutant emissions. According to UNFCCC/MMR reporting obligations split between national and memo international aviation emissions is not required. To find a way for separately reporting emissions outside of national totals would take to much effort with no significant improvement of national totals. According to 2018 in-depth EU NECD review explanation of the progress is included. We tried to obtain separate data, but we were not successful. Statistical Office, which is our main fuel data provider, does not provide those data. Since we have no possibilities for obtain reliable data, we are not planning to report data separately in next annual submission.

### **Domestic aviation cruise (civil): NFR Code 1A3aii(ii)-memo items**

Notation Key "IE" (included elsewhere) was used for Domestic aviation cruise (civil), since all fuel used for this sector was reported under 1A3aii(i) Domestic aviation LTO (civil). Overestimation of national emissions due to inclusion of memo category 1A3aii(ii) to national totals is below 1 % and does not have a significant impact on national inventory. Since this source is of less importance Tier 1 method was used for emission calculation. To much effort with no significant improvement of national totals would be needed for separate reporting of 1A3aii(ii) emissions outside of national totals. According to 2018 in-depth EU NECD review explanation of the progress is included. We tried to obtain separate data, but we were not successful. Statistical Office, which is our main fuel data provider, does not provide those data. Since we have no possibilities for obtain reliable data, we are not planning to report data separately in next annual submission.

### 3.4 Small Combustion and Non-road mobile sources and machinery (1. A. 4)

This chapter covers the methods and data needed to estimate stationary combustion emissions in smaller-scale combustion units than those in Chapter 1A1, Energy industries. The small combustion installations included in this chapter are mainly intended for heating and provision of hot water in residential and commercials/institutional sectors.

This chapter also provides the estimation of combustion emissions from non-road mobile sources and machinery. It covers a mixture of 'other' equipment which is distributed across a wide range of industry sectors. All the equipment covered uses reciprocating engines, fuelled with liquid hydrocarbon-based fuels. They comprise both diesel and petrol engined machinery.

This category is very important source of air pollutant emissions. It mostly contributes to total emissions of particulate matter, CO, PAHs, dioxins/furans. It is important source of Cd, Cr, Ni, Zn, NMVOC, NO<sub>x</sub>, HCB as well. The most important source of these pollutants is residential sector, mostly due to much of biomass burning.

Sectors covered in this chapter are:

NFR Codes:

1A4ai	Commercial/institutional: Stationary
1A4bi	Residential: Stationary
1A2gvii	Mobile Combustion in manufacturing industries and construction
1A4cii	Agriculture/Forestry/Fishing: Off-road vehicles and other machinery
1A3ei	Pipeline transport

#### 3.4.1 Commercial/institutional: Stationary (NFR Code 1A4ai) and Residential: Stationary (NFR Code 1A4bi)

##### Introduction

The small combustion installations included in this chapter are mainly intended for heating and provision of hot water in residential and commercials/institutional sectors. Some of these installations are also used for cooking, primarily in the residential sector. Emissions from smaller combustion installations are significant due to their numbers, different type of combustion techniques employed, and range of efficiencies and emissions.

##### Methodology

To estimate emissions from combustion in manufacturing industries and construction the following formulas have been used:

$$E = m \times \text{NCV} \times \text{EF}$$

*Equation 1*

E - emission (g)

m - quantity of fuel combusted (t)

NCV - net calorific value (TJ/kt)

EF - emission factor per energy of fuel (g/GJ)

$$E = m \times EF$$

Equation 2

E - emission (g)

m - quantity of fuel combusted (t)

EF - emission factor per quantity of fuel (g/t)

To estimate SO<sub>x</sub> emissions in some cases the following two equations for calculation of EF were used:

$$EF_{SO_x} = [S] \times 20000 / NCV$$

Equation 3

EF<sub>SO<sub>x</sub></sub> - SO<sub>x</sub> emission factor (g/GJ)

[S] - sulphur content of the fuel (% w/w)

NCV - net calorific value (GJ/t)

2 - ratio of the relative molecular mass of SO<sub>x</sub> to sulphur

$$EF_{SO_x} = [S] \times 19000 / NCV$$

Equation 4

EF<sub>SO<sub>x</sub></sub> - SO<sub>x</sub> emission factor (g/GJ)

[S] - sulphur content of the fuel (% w/w)

NCV - net calorific value (GJ/t)

1,9 - ratio of the relative molecular mass of SO<sub>x</sub> to sulphur, considering 5 % absorption in the ash

### Activity data

Data on the consumption of fuels in the commercial sector and households were obtained from Statistical Office of the Republic of Slovenia (SORS). Lignite, domestic and imported sub-bituminous coal, heavy fuel oil, residual fuel oil, LPG, natural gas, wood and other biomass have been used in both categories. Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.16: Fuel used in the Other sectors 1980–2017).

### Net calorific values

Net calorific values have been taken from SORS. The values for solid fuel varies from year to year but for the liquid and gaseous fuel almost the same values have been used for the entire period, as these types of fuel do not change a lot from year to year.

Table 3.4.1.1 NCVs for the fuel used in commercial and residential sector

Year	Lignite – domestic	Sub-bituminous Coal - domestic	Sub-bituminous Coal - imported	Residual Fuel Oil	Heavy Fuel Oil	LPG	Natural Gas	Wood and Other Biomass
	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/kt	TJ/Mm3	TJ/kt
1980	9,360	12,980		41,800	39,700	46,050	33,500	12,170
1981	9,330	11,570		41,800	39,700	46,050	34,100	12,170
1982	9,330	11,570		41,900	39,800	46,000	33,490	12,170
1983	9,610	11,180		41,900	39,800	46,000	33,800	12,170
1984	9,590	11,420		41,900	40,000	46,000	33,500	12,170
1985	9,430	11,690		41,900	39,800	46,050	33,500	12,170
1986	9,390	12,850		41,820	39,740	46,000	33,500	12,170
1987	9,650	11,820		41,780	39,800	46,000	33,500	12,170
1988	9,440	12,000		41,710	39,800	46,000	34,080	12,170

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1989	9,820	12,050		41,850	39,800	46,000	34,100	12,170
1990	9,810	12,760		41,870	39,800	46,000	34,100	12,170
1991	9,980	12,879		41,880	39,800	46,000	34,100	12,170
1992	10,260	12,589		41,900	39,900	46,000	34,100	12,170
1993	10,070	13,351		41,900	39,800	46,000	34,100	12,170
1994	9,960	12,666		41,900	39,860	46,000	34,100	12,170
1995	10,220	17,404		41,900	40,000	46,000	34,100	12,170
1996	9,690	16,353		41,900	40,000	46,000	34,100	12,170
1997	9,610	18,203		41,900	40,000	46,050	34,080	12,170
1998	10,010	18,531		41,900	40,000	46,050	34,080	12,170
1999	9,690	18,563		41,900	40,000	46,050	34,080	12,170
2000	10,170	17,983		41,900	40,000	46,050	34,080	12,261
2001	10,660	16,353		41,900	40,000	46,050	34,080	12,511
2002	10,350	19,000		41,900	40,000	46,050	34,080	12,766
2003	10,138	19,000		41,900	40,000	46,050	34,080	13,027
2004	10,138	19,000		41,900		46,050	34,080	13,293
2005	10,803		17,000	42,600		46,050	34,080	13,564
2006			17,318	41,900		46,050	34,072	13,841
2007			16,863	42,600		46,050	34,076	14,123
2008			16,407	42,600		46,050	34,096	14,412
2009			15,952	42,600		46,050	34,080	14,742
2010			16,155	42,600		46,050	34,080	14,747
2011			15,985	42,600		46,050	34,087	14,777
2012			16,032	42,600		46,050	34,093	14,799
2013			16,457	42,600		46,050	34,079	14,805
2014			15,734	42,600		46,050	34,083	14,809
2015			16,360	42,600		46,050	34,086	14,813
2016			16,575	42,600		46,050	34,087	14,816
2017			16,000	42,600		46,050	34,085	14,821

### Emission factors

For calculating emissions of individual gases in commercial and residential sector following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

Table 3.4.1.2 Emission factors used for domestic and imported sub-bituminous coal and lignite in residential sector for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	110	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
SO <sub>x</sub>	Equation 4	[S] (% w/w) See Table 3.2.1.10	Slovene national legislation relating quality of liquid fuels
CO	4600	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
NM VOC	484	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
NH <sub>x3</sub>	0,3	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
PM <sub>10</sub>	404	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
PM <sub>2.5</sub>	398	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
TSP	444	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
BC	25,472	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36

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<b>Cd</b>	1,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Pb</b>	130	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Hg</b>	5,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>As</b>	2,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Cr</b>	11,2	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Cu</b>	22,3	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Ni</b>	12,7	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Se</b>	120	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Zn</b>	220	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Dioxins/ Furans</b>	800	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Benzo(a)pyrene</b>	230	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Benzo(b)fluoranthene</b>	330	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Benzo(k)fluoranthene</b>	130	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>Indeno(1,2,3-cd)pyrene</b>	110	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>HCB</b>	0,62	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36
<b>PCB</b>	170	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-3, pg 36

Table 3.4.1.3 Emission factors used for residual fuel oil in residential sector for 1980 - 2017

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	51	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>SO<sub>x</sub></b>	<i>Equation 3</i>	[S] (% w/w) See Table 3.2.1.12	Slovene national legislation relating quality of liquid fuels
<b>CO</b>	57	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>NM VOC</b>	0,69	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>PM<sub>10</sub></b>	1,9	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>PM<sub>2.5</sub></b>	1,9	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>TSP</b>	1,9	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>BC</b>	0,162	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Cd</b>	0,001	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Pb</b>	0,012	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Hg</b>	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>As</b>	0,002	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38

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<b>Cr</b>	0,2	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Cu</b>	0,13	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Ni</b>	0,005	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Se</b>	0,002	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Zn</b>	0,42	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Dioxins/ Furans</b>	5,9	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Benzo(a)pyrene</b>	80	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Benzo(b)fluoranthene</b>	40	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Benzo(k)fluoranthene</b>	70	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38
<b>Indeno(1,2,3-cd)pyrene</b>	160	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-5, pg 38

**Table 3.4.1.4 Emission factors used for natural gas and liquefied petroleum gas oil in residential sector for 1980 - 2017**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	51	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>CO</b>	26	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>SO<sub>x</sub></b>	0,3	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>NMVOC</b>	1,9	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>PM<sub>10</sub></b>	1,2	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>PM<sub>2.5</sub></b>	1,2	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>TSP</b>	1,2	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>BC</b>	0,0648	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Cd</b>	0,00025	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Pb</b>	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Hg</b>	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>As</b>	0,12	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Cr</b>	0,00076	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Cu</b>	0,000076	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Ni</b>	0,00051	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Se</b>	0,011	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Zn</b>	0,0015	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Benzo(a)pyrene</b>	0,56	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37

<b>Benzo(b)fluoranthene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Benzo(k)fluoranthene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Indeno(1,2,3-cd)pyrene</b>	0,84	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37
<b>Dioxins/ Furans</b>	1,5	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-4, pg 37

**Table 3.4.1.5 Emission factors used for wood and other biomass in residential sector for 1980 - 2017**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>SO<sub>x</sub></b>	11	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Cd</b>	13	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Pb</b>	27	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Hg</b>	0,56	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>As</b>	0,19	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Cr</b>	23	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Cu</b>	6	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Ni</b>	2	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Se</b>	0,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>Zn</b>	512	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39
<b>HCB</b>	5	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-6, pg 39

For calculation of NO<sub>x</sub>, CO, NH<sub>3</sub>, NMVOC, PCB, dioxins/furans, PAHs and particulate matter emissions from wood combustion in residential plants Tier 2 emission factors were used. We have estimated a share of different types of technologies for wood combustion in residential sector for the period 2005 - 2017. 2005 data was applied for the period 1980 - 2004 since no data on structure of heating equipment in residential sector is available prior 2005.

In the year 2017 there were 67 % conventional boilers < 50 kW burning wood and similar wood waste, 13 % advanced / ecolabelled stoves and boilers burning wood, 4 % pellet stoves and boilers burning wood pellets, 1 % open fireplaces burning wood, 15 % conventional stoves burning wood and similar wood waste.

Emission factors have been obtained from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Small combustion, Table 3-39, pg. 82, Table 3-40, pg. 84, Table 3-43, pg. 88, Table 3-42, pg. 87, Table 3-44, pg. 90.

**Table 3.4.1.6 Emission factors used for wood and other biomass in residential sector for NO<sub>x</sub>, NH<sub>3</sub>, NMVOC, CO, PM<sub>10</sub>, PM<sub>2.5</sub> and TSP and BC**

Year	NMVOC	NH <sub>3</sub>	NO <sub>x</sub>	CO	PM <sub>2.5</sub>	PM <sub>10</sub>	TSP	BC
Unit	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ	g/GJ
<b>2005 and before</b>	382	70,8	75,8	3857	491	503	525	72
<b>2006</b>	381	70,3	76,0	3830	487	498	520	71
<b>2007</b>	374	70,4	76,7	3832	480	491	513	71
<b>2008</b>	375	69,6	76,6	3786	476	487	509	70
<b>2009</b>	374	68,9	76,6	3748	471	482	503	69
<b>2010</b>	373	68,2	76,6	3711	467	478	499	68
<b>2011</b>	367	67,7	76,9	3680	461	471	492	68
<b>2012</b>	364	67,2	77,1	3647	456	466	487	67
<b>2013</b>	362	66,9	77,2	3629	453	463	484	67
<b>2014</b>	365	66,2	76,5	3599	455	466	487	66
<b>2015</b>	358	65,6	77,2	3559	445	455	475	65
<b>2016</b>	355	65,2	77,4	3534	439	449	469	65
<b>2017</b>	353	64,5	77,4	3498	436	446	466	64

**Table 3.4.1.7 Emission factors used for wood and other biomass in residential sector for PCB, dioxins/furans, PAHs**

Year	PCB	Dioxins/ Furans	Benzo(a) pyrene	Benzo(b) fluoranthene	Benzo(k) fluoranthene	Indeno(1,2,3-cd) pyrene
Unit	microg/GJ	ng/GJ	mg/GJ	mg/GJ	mg/GJ	mg/GJ
<b>1990-2005</b>	0,0568	563	114,3	51,7	62,3	15,7
<b>2006</b>	0,0561	558	112,8	51,5	61,4	15,8
<b>2007</b>	0,0562	552	112,9	49,7	62,1	14,1
<b>2008</b>	0,0551	546	110,6	50,1	60,4	15,0
<b>2009</b>	0,0542	540	108,8	50,0	59,2	15,3
<b>2010</b>	0,0535	535	107,2	50,0	58,1	15,6
<b>2011</b>	0,0530	528	106,2	49,1	57,8	15,0
<b>2012</b>	0,0525	523	105,1	48,7	57,1	14,8
<b>2013</b>	0,0522	520	104,5	48,3	56,9	14,6
<b>2014</b>	0,0517	521	103,2	49,7	55,5	16,2
<b>2015</b>	0,0509	510	101,6	48,1	54,9	15,1



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2016	0,0504	504	100,6	47,5	54,6	14,6
2017	0,0498	500	99,2	47,5	53,6	14,9

It is unclear for solid fuels, gaseous fuel and liquid fuels whether emission factors represent filterable PM emissions or total (filterable and condensable) emissions. TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors for biomass represent total particulate emissions (filterable and condensable) emissions.

**Table 3.4.1.8 Emission factors used for domestic sub-bituminous coal and lignite in commercial sector for 1980 - 2004**

Pollutant	Value	Unit	References
NO <sub>x</sub>	173	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
SO <sub>x</sub>	Equation 4	[S] (% w/w) See Table 3.2.1.10	Slovene national legislation relating quality of liquid fuels
CO	931	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
NM VOC	88,8	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
PM <sub>10</sub>	117	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
PM <sub>2.5</sub>	108	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
TSP	124	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
BC	6,912	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Cd	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Pb	134	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Hg	7,9	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
As	4	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Cr	13,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Cu	17,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Ni	13	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Se	1,8	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Zn	200	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Dioxins/ Furans	203	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Benzo(a)pyrene	45,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Benzo(b)fluoranthene	58,9	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Benzo(k)fluoranthene	23,7	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
Indeno(1,2,3-cd)pyrene	18,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40

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<b>HCB</b>	0,62	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40
<b>PCB</b>	170	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-7, pg 40

**Table 3.4.1.9 Emission factors used for heavy fuel oil and residual fuel oil in commercial sector for 1980 - 2017**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>NO<sub>x</sub></b>	306	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>SO<sub>x</sub></b>	<i>Equation 3</i>	[S] (% w/w) See Table 3.2.1.12	Slovene national legislation relating quality of liquid fuels
<b>CO</b>	93	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>NMVOC</b>	20	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>PM<sub>10</sub></b>	20	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>PM<sub>2.5</sub></b>	18	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>TSP</b>	21	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>BC</b>	10,08	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Cd</b>	0,15	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Pb</b>	8	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Hg</b>	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>As</b>	0,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Cr</b>	10	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Cu</b>	3	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Ni</b>	125	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Se</b>	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Zn</b>	18	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Dioxins/ Furans</b>	6	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Benzo(a)pyrene</b>	1,9	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Benzo(b)fluoranthene</b>	15	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Benzo(k)fluoranthene</b>	1,7	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>Indeno(1,2,3-cd)pyrene</b>	1,5	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>HCB</b>	0,22	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42
<b>PCB</b>	0,13	ng/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-9, pg 42

Table 3.4.1.10 Emission factors used for natural gas, liquefied petroleum gas and gaseous biomass in commercial sector for 1980 - 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	74	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
CO	29	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
SO <sub>x</sub>	0,67	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
NM VOC	23	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
PM <sub>10</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
PM <sub>2.5</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
TSP	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
BC	0,0312	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cd	0,0009	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Pb	0,011	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Hg	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
As	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cr	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cu	0,0026	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Ni	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Se	0,058	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Zn	0,73	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(a)pyrene	0,72	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(b)fluoranthene	2,9	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(k)fluoranthene	1,1	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Indeno(1,2,3-cd)pyrene	1,08	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Dioxins/ Furans	0,52	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41

Table 3.4.1.11 Emission factors used for wood in commercial sector for 1980 - 2005

Pollutant	Value	Unit	References
NO <sub>x</sub>	91	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
CO	570	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
NM VOC	300	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
NH <sub>3</sub>	37	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
SO <sub>x</sub>	11	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43

<b>PM<sub>10</sub></b>	143	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>PM<sub>2.5</sub></b>	140	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>TSP</b>	150	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>BC</b>	39,2	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Cd</b>	13	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Pb</b>	27	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Hg</b>	0,56	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>As</b>	0,19	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Cr</b>	23	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Cu</b>	6	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Ni</b>	2	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Se</b>	0,5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Zn</b>	512	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Benzo(a)pyrene</b>	10	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Benzo(b)fluoranthene</b>	16	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Benzo(k)fluoranthene</b>	5	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Indeno(1,2,3-cd)pyrene</b>	4	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>Dioxins/ Furans</b>	100	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>PCB</b>	0,06	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43
<b>HCB</b>	5	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-10, pg 43

It is unclear for solid fuels, gaseous fuel and liquid fuels whether emission factors represent filterable PM emissions or total (filterable and condensable) emissions.

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors for solid biomass represent filterable emissions.

## Emissions

These two subsectors are very important source of CO, particulate matter, dioxins/furans, PAHs and Zn, Cd, Cr. In 2017 these two sectors contributed 65 % of CO emissions, 61 to 73 % of various particulate matter, 62 % of dioxins/furans and 79 % of PAHs national emissions. Emissions of CO, PAHs, dioxins/furans have decreased from 1990 to 2017 due to shift in the fuel mix from solid fuels to natural gas. But distinctive increase of all emissions, including particulate matter, was observed in 2008 due to higher use of wood biomass in residential sector. This was a result of economic crisis and high price of petroleum products as well as state measures to promote renewable energy sources.

## Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Data on wood consumption in 1A4bi Residential: Stationary for the period 1990-2016 has been improved and related emissions have been recalculated for that period.

### **Category-Specific QA/QC and Verification**

According to 2018 in-depth EU NECD review information on biomass activity data used in Commercial/institutional: Stationary subsector was included. This sector comprises solid biomass for the period 1980-2005 and biogas fuels for the period 2008-2017. In the 2017 submission the same amount of wood consumption have been used for the period 1990-2000. For 2018 submission we have used improved data from the SORS for the year 1990 and 2000 while the amount of wood used in the period 1991 – 1999 was interpolated. Since 2000 the data in the inventory and SORS data are the same. In the commercial/institutional sector since 2006 no wood biomass has been consumed any more. Biomass, which has been used since 2008, is biogas. Data in Annex 1 was checked and corrected.

### **Future Improvements**

No improvements are planned for next submission.

## **3.4.2 Mobile Combustion in manufacturing industries and construction**

NFR Code 1A2gvii

### **Introduction**

This sector includes emissions from construction land-based mobile machinery. Different types of vehicles and machinery are used in building industry (asphalt and concrete pavers, roller, cement and mortar mixers...). Emissions originate from the combustion of fuel (diesel and gasoline) to power this equipment. Contribution of emissions to the total national inventory is of less importance. Contribution of NO<sub>x</sub> emissions is 2 % and black carbon 1,6 %, other pollutants contributed less than 1 % in 2017.

### **Methodology**

To estimate exhaust emissions from off-road construction equipment the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (kg/t)

### **Activity data**

Data on amount of diesel and gasoline used for non-road mobile machinery in construction sector were obtained from SORS. Data are available for the period 1986-2017. Amount of diesel combusted has been much bigger than gasoline used. 25906 t of diesel and 191 t of gasoline

were consumed in the year 2017. Fuel consumption for the whole period is shown in the Annex 1 to the IIR (Table 1.12: Fuel Consumption in Mobile Combustion in manufacturing industries and construction).

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

**Table 3.4.2.1 Emission factors for leaded and unleaded gasoline used in construction**

Pollutant	Value	Unit	References
NO <sub>x</sub>	7,117	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
SO <sub>x</sub>	Values used for road transport (Table 3.3.1.2)	kg/t	Slovene national legislation relating quality of liquid fuels
CO	770,368	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
NM VOC	18,893	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
NH <sub>3</sub>	0,004	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
PM <sub>10</sub>	0,157	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
PM <sub>2.5</sub>	0,157	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
TSP	0,157	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
BC	0,008	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Cd	0,010	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Cu	1,7	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Cr	0,05	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Ni	0,07	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Se	0,01	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Zn	1	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Pb (Unleaded gasoline)	0,033	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-10
Pb (Leaded gasoline)	200	g/t	Slovene national legislation relating quality of liquid fuels
Benzo(a)pyrene	0,0400	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
Benzo(b)fluoranthene	0,0400	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
Benzo(k)fluoranthene	0,0039	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
Indeno(1,2,3-cd)pyrene	0,0089	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8

Table 3.4.2.2 Emission factors for diesel used in construction

Pollutant	Value	Unit	References
NO <sub>x</sub>	32,629	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
SO <sub>x</sub>	Values used for road transport (Table 3.3.1.2)	kg/t	Slovene national legislation relating quality of liquid fuels
CO	10,774	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
NM VOC	3,377	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
NH <sub>3</sub>	0,008	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
PM <sub>10</sub>	2,104	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
PM <sub>2.5</sub>	2,104	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
TSP	2,104	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
BC	1,304	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Cd	0,0100	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Cu	1,7	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Cr	0,05	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Ni	0,07	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Se	0,01	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Zn	1	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Benzo(a)pyrene	0,0300	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Benzo(b)fluoranthene	0,0500	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
Benzo(k)fluoranthene	0,0344	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
Indeno(1,2,3-cd)pyrene	0,0079	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors represent total PM emissions (filterable and condensable fractions).

### Emissions

In the period 2006-2008 the highest liquid fuel consumption was observed with the peak in the year 2006. This increase is associated with the economic situation in Slovenia at that time. A high economic growth in the period 2004-2008 had influenced the increase of investments into real estates. According to the SORS data, the highest number of building permits have been issued just in 2006 what means that more fuel demanding phases in construction of buildings (excavation of construction pits) had happened in 2006. The construction of highways has been also rapidly expanding in this period.

### Category-Specific QA/QC and Verification

According to 2017 in-depth EU NECD review recommendations new EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for emissions calculation. Data on fuel consumption have been checked and compare with the SORS data. No inconsistencies have been found.

### Recalculations

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Future Improvements

No improvements are planned for next submission.

## 3.4.3 Agriculture/Forestry/Fishing: Off-road vehicles and other machinery

NFR Code 1A4cii

### Introduction

This sector includes emissions resulting from consumption of fuel used for off-road vehicles and other machinery in agriculture and forestry land based mobile machinery. Fishing is excluded from this sector and is reported under 1A3b Road transport. Exhaust emissions from non-road mobile machinery arisen from the combustion of diesel and gasoline in agriculture and forestry. Emissions of NO<sub>x</sub>, NMVOC, CO and particulate matter contribute a few percent to the total national emissions. Contributions of other pollutants are below 1 %.

### Methodology

To estimate exhaust emissions from off-road vehicles and other machinery used in agriculture and forestry the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (kg/t)

### Activity data

The consumption of fuels until year 2000 has been calculated from data on fuel consumption in state owned agriculture enterprises and corresponding agriculture land. Data were obtained from SORS. The same energy intensity have been used to calculate fuel used on total agricultural land. For estimation of fuel consumption in agriculture from year 2000 onwards, we used the same energy intensity (fuel consumption/ha of land) as observed in 2000.

The consumption of fuels in the entire forestry is estimated on the basis of consumption of fuel



in state-owned logging enterprises. For the state-owned sector, data are available for the consumption of fuel and cut, for private sector only data on cut. First, the consumption per m<sup>3</sup> of cut in state owned logging enterprises is estimated. Based on these estimates and data on total cut, the estimate of consumption in the whole of forestry is calculated. Before 2005 there were no separate data on consumption of gasoline and gas, only the total consumption. Consequently, the split is done considering the split in agriculture (10 % gasoline, 90 % gas oil), presuming that the same amount of fuels is consumed per m<sup>3</sup> of felled wood in private forestry as in state forestry. For the period 2005 - 2017 we have obtained direct data on amount of fuel used in forestry from SORS.

Fuel consumption in agriculture and forestry for the whole period is shown in the Annex 1 to the IIR (Table 1.13: Fuel Consumption in Agriculture/Forestry/Fishing: Off-road vehicles and other machinery).

### Emission factors

In calculating emissions of individual gases, following emission factors from new EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

**Table 3.4.3.1 Emission factors for leaded and unleaded gasoline used in agriculture and forestry**

Pollutant	Value	Unit	References
NO <sub>x</sub>	2,765	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
SO <sub>x</sub>	Values used for road transport (Table 3.3.1.2)	kg/t	Slovene national legislation relating quality of liquid fuels
CO	620,793	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
NM VOC	227,289	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
NH <sub>3</sub>	0,003	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
PM <sub>10</sub>	3,762	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
PM <sub>2.5</sub>	3,762	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
TSP	3,762	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
BC	0,188	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Cd	0,010	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Pb (Unleaded gasoline)	0,033	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 24, Table 3-10
Pb (Leaded gasoline)	200	g/t	Slovene national legislation relating quality of liquid fuels
Cu	1,7	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Cr	0,05	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 25
Ni	0,07	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
Se	0,01	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
Zn	1	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26

<b>Benzo(a)pyrene</b>	0,04	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
<b>Benzo(b)fluoranthene</b>	0,04	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 26
<b>Benzo(k)fluoranthene</b>	0,0039	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8
<b>Indeno(1,2,3-cd)pyrene</b>	0,0089	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, pg 23, Table 3-8

Table 3.4.3.2 Emission factors for diesel used in agriculture and forestry

Pollutant	Value	Unit	References
<b>SO<sub>x</sub></b>	Values used for road transport (Table 3.3.1.2)	kg/t	Slovene national legislation relating quality of liquid fuels
<b>CO</b>	11,469	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 23
<b>NH<sub>3</sub></b>	0,008	kg/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 23
<b>Cd</b>	0,010	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Cu</b>	1,7	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Cr</b>	0,05	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Ni</b>	0,07	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Se</b>	0,01	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Zn</b>	1	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Benzo(a)pyrene</b>	0,030	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Benzo(b)fluoranthene</b>	0,050	g/t	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-1, pg 24
<b>Benzo(k)fluoranthene</b>	0,0344	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, Table 3-8, pg 23
<b>Indeno(1,2,3-cd)pyrene</b>	0,0079	g/t	Emission Inventory Guidebook, 2016, Exhaust emissions from road transport, Table 3-8, pg 23

For calculation of NO<sub>x</sub>, NMVOC and particulate matter emissions from diesel machinery in agriculture and forestry Tier 3 emission factors were used. Vehicles population, predominantly tractors is split into different types, ages and power ranges. The baseline emission factors for regulated diesel engines and machinery are taken as the EU type approval values (expressed in g/kWh). Shares of tractors with different age, power range and technology were taken into consideration for emission calculation.

Table 3.4.3.3 Emission factors for NMVOC for diesel used in agriculture and forestry for 1990-2017

Year	NMVOC	Unit	References
<b>1990-2005</b>	246	g/GJ	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-6, pg 38 and expert evaluation
<b>2006</b>	243	g/GJ	

2007	233	g/GJ	
2008	222	g/GJ	
2009	214	g/GJ	
2010	209	g/GJ	
2011	205	g/GJ	
2012	199	g/GJ	
2013	193	g/GJ	
2014	188	g/GJ	
2015	182	g/GJ	
2016	178	g/GJ	
2017	162	g/GJ	

Table 3.4.3.4 Emission factors for NO<sub>x</sub> for diesel used in agriculture and forestry for 1980-2017

Year	NO <sub>x</sub>	Unit	References
1980-2005	1237	g/GJ	
2006	1220	g/GJ	
2007	1182	g/GJ	
2008	1122	g/GJ	
2009	1083	g/GJ	
2010	1057	g/GJ	
2011	1031	g/GJ	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-6, pg 38 and expert evaluation
2012	1010	g/GJ	
2013	986	g/GJ	
2014	960	g/GJ	
2015	926	g/GJ	
2016	908	g/GJ	
2017	891	g/GJ	

Table 3.4.3.5 Emission factors for PM<sub>10</sub>, PM<sub>2.5</sub>, TSP and BC for diesel used in agriculture and forestry for 2000-2017

Year	PM <sub>2.5</sub>	PM <sub>10</sub>	TSP	BC	Unit	References
2000-2005	99	105	110	62	g/GJ	Emission Inventory Guidebook, 2016, Non-road mobile sources and machinery, Table 3-6, pg 38 and

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2006	97	103	108	60	g/GJ	expert evaluation
2007	93	98	103	58	g/GJ	
2008	88	93	98	55	g/GJ	
2009	85	90	95	53	g/GJ	
2010	83	88	92	52	g/GJ	
2011	81	86	90	50	g/GJ	
2012	79	83	88	49	g/GJ	
2013	76	81	85	47	g/GJ	
2014	74	78	82	46	g/GJ	
2015	71	75	79	44	g/GJ	
2016	69	73	77	43	g/GJ	
2017	68	72	75	42	g/GJ	

TSP, PM<sub>10</sub>, PM<sub>2.5</sub> emission factors represent total PM emissions (filterable and condensable fractions).

### Recalculations

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-Specific QA/QC and Verification

According to 2017 and 2018 in-depth EU NECD review recommendations we performed an examination of gasoline-powered equipment used in agriculture and forestry. According to logging companies all gasoline used in forestry is applied in two-stroke chain saws. No four-stroke equipment is used. Due to economical reasons all other machinery is diesel - powered. We put additional effort to obtain reliable information on use of gasoline equipment in forestry. More sources were checked, including Statistical Office of Republic of Slovenia. No data is available on four-stroke gasoline in forestry.

Examination of gasoline-powered equipment used in agriculture was performed as well. More sources were checked, including Statistical Office of Republic of Slovenia. We did not get any better and reliable information on gasoline powered agriculture equipment. Since gasoline contributes only a very small part (7 %) to total fuel consumption and we do not have any precise and reliable data, we decided to use Tier 1 emission factors for gasoline equipment. Tier 3 EFs are applied for emissions from diesel-powered equipment, whereas Tier 1 default EF are applied for two-stroke gasoline equipment.

### Future Improvements

No improvement is planned for next submission.

### 3.4.4 Pipeline transport

NFR Code 1A3ei

#### Introduction

This category includes emissions from natural gas combusted on compressor station. Emissions from this source are negligible. They are far below 0,01 %.

#### Methodology

To estimate emissions the following methodology has been adopted.

$$E = m \times \text{NCV} \times \text{EF}$$

E – emission (mg)  
 m – quantity of fuel combusted (m<sup>3</sup>)  
 EF – emission factor per energy of fuel (g/GJ)  
 NCV - net calorific value (MJ/m<sup>3</sup>)

#### Activity data

We have obtained data on natural gas used on compressor station from the company which is the owner of this compressor station. The data are available from 2008. Activity data for 2017 is 2429587 m<sup>3</sup> of natural gas.

#### Net calorific values

Net calorific values have been taken from SORS.

**Table 3.4.4.1 NCVs for natural gas used on compressor station**

Year	Natural Gas
	MJ/m <sup>3</sup>
2008	34,096
2009	34,080
2010	34,080
2011	34,087
2012	34,093
2013	34,079
2014	34,083
2015	34,086
2016	34,087
2017	34,085

#### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

Table 3.4.4.2 Emission factors used for natural gas on compressor station for 2008 – 2017

Pollutant	Value	Unit	References
NO <sub>x</sub>	74	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
CO	29	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
NM VOC	23	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
SO <sub>x</sub>	0,67	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
PM <sub>10</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
PM <sub>2.5</sub>	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
TSP	0,78	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
BC	0,0312	g/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cd	0,0009	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Pb	0,011	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Hg	0,54	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
As	0,1	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cr	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Cu	0,0026	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Ni	0,013	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Se	0,058	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Zn	0,73	mg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(a)pyrene	0,72	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(b)fluoranthene	2,9	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Benzo(k)fluoranthene	1,1	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Indeno(1,2,3-cd)pyrene	1,08	microg/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41
Dioxins/ Furans	0,52	ng I-TEQ/GJ	Emission Inventory Guidebook, 2016, Small combustion, Table 3-8, pg 41

It is unclear whether emission factors for TSP, PM<sub>10</sub>, PM<sub>2.5</sub> represent filterable PM emissions or total (filterable and condensable) emissions.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2008-2017.

### Category-Specific QA/QC and Verification

According to 2017 in-depth EU NECD review recommendations emission factors and activity

data was checked. New EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for emissions calculation. Notation keys were revised as well. "NE" was applied for NH<sub>3</sub> emissions.

**Future Improvements**

No improvement is planned for this category.

**Commercial / institutional: Mobile: NFR Code 1A4aii**

Fuel used for commercial and institutional land-based mobile machinery is included 1A3b Road transport. Notation Key "IE" (included elsewhere) was therefore used for this sector.

**Residential: Household and gardening (mobile): NFR Code 1A4bii**

Equipment used in household and gardening are fuelled at regular petrol stations. Separation of fuel sold to road vehicles and household and gardening landbased mobile machinery is not possible. There is no data on fuel used for mobile sources in household and gardening and Statistical office has no intension to collect this data in the near future. We believe that amount of fuel used for this purpose is very small. Notation Key "IE" (included elsewhere) was used for this sector since all fuel used for household and gardening was reported under 1A3b Road transport. We tried to obtain separate data, but we were not successful. Statistical Office, which is our main fuel data provider, does not provide those data. Since we have no possibilities for obtain reliable data, we are not planning to report data separately in next annual submission.

**Agriculture/Forestry/Fishing: Stationary NFR Code 1A4ci**

Fuel used in stationary agriculture and forestry installations is included under 1A4bi Residential: Stationary. Notation Key "IE" (included elsewhere) was therefore used for this sector.

**Agriculture/Forestry/Fishing: National fishing: NFR Code 1A4ciii**

Emissions from fishing are not included in this sector, because the data on the fuel used for this purpose are not available separately. According to The European Community Fishing Fleet Register, there have been only 175 active fishing motor vessels in Slovenia at the end of 2016. Majority of them (150) are less than 10 m long and the longest boat has only 18 m. Due to the unresolved see border with Croatia and due to the EU legislation on Common Fisheries Policy (the subsidies are given to fishermen if they give up fishing and destroy the vessels) the number of vessels and fishermen are decreasing from year to year. Fuel used for fishing vessels has been sold on four petrol stations at Adriatic coast (Izola Pier, Lucija Pier, Marina Koper and Piran Pier). These patrol stations have been selling fuel to road vehicles as well. Separation of fuel sold to road vehicles and fishing vessels is not possible. Notation Key "IE" (included elsewhere) was used for fishing, since all fuel used for this sector was reported under 1A3b Road transport. We tried to obtain separate data, but we were not successful. Statistical Office, which is our main fuel data provider, does not provide those data. Since we have no possibilities for obtain reliable data, we are not planning to report data separately in next annual submission.

**Other stationary (including military): NFR Code 1A5a**

Fuel used in other small stationary installations is included in 1A4ai Commercial/institutional: Stationary. Notation Key "IE" (included elsewhere) was therefore used for this sector.

**Other (please specify in the IIR): NFR Code 1A3eii**

Notation Key "NO" (not occurring) was used for this sector, since there is no other additional emissions in Slovenia.



### 3.5 Fugitive emissions from fuels (1. B)

This chapter covers fugitive emissions from solid fuels and oil and natural gas.

Sectors covered in this chapter are:

NFR Codes:

1B1a	Fugitive emissions from solid fuels: Coal mining and handling
1B2ai	Fugitive emissions oil: Exploration, production, transport
1B2aiv	Fugitive emissions oil: Refining / storage
1B2av	Distribution of oil products
1B2b	Fugitive emissions from natural gas (exploration, production, processing, transmission, storage, distribution and other)
1B2c	Venting and flaring (oil, gas, combined oil and gas)

#### 3.5.1 Fugitive emissions from solid fuels: Coal mining and handling

NFR Code 1B1a

##### Introduction

This chapter encompasses emissions arising from the production, processing, and storage of coal from underground coal mines. The extraction and treatment of coal result mainly in emissions of greenhouse gas methane. The most important component of those emissions is CH<sub>4</sub> emissions that arise in mining and post-mining activities although CO<sub>2</sub> emissions occur as well. However, also non-methane volatile organic compounds and particulate matter are emitted. Emissions of NMVOC have been calculated for the period 1990-2017, emissions of particulate matter for the period 2000-2017. Emissions of NMVOC and particulate matter from this source contributed in 2017 a few percent to total national emissions.

##### Methodology

To estimate fugitive emissions from coal mining and handling the following methodology has been adopted:

$$E = m \times EF$$

E – emission (g)

m – quantity of fuel combusted (t)

EF – emission factor per quantity of fuel (g/t)

##### Activity data

Data on excavated quantities of coal according to individual coalmines are obtained from Statistical Office of the Republic of Slovenia (SORS). Only one coal mine has been in operations in Slovenia in the year 2017. Data on excavated quantities of coal according to individual coalmines are presented on the Table 3.5.1.1.

**Table 3.5.1.1 Excavation of coal in Slovenia 1986 – 2017**

Pit	1986	1990	2000	2005	2010	2013	2015	2016	2017	Closed in
Unit	kt	kt	kt	kt	kt	kt	kt	kt	kt	
Velenje	5001	4210	3743	3945	4011	3826	3168	3349	3356	
Trbovlje - Hrastnik	1242	905	737	594	419	51				2013
Zagorje	315	244								1997
Senovo	120	108								1996
Kanižarica	126	94								1996
Laško	25									1990
<b>Total Coal Excavation</b>	<b>6828</b>	<b>5561</b>	<b>4480</b>	<b>4540</b>	<b>4430</b>	<b>4278</b>	<b>3168</b>	<b>3349</b>	<b>3356</b>	

### Emission factors

Emission factors for PM<sub>2.5</sub>, PM<sub>10</sub> and TSP were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Fugitive emissions, Fugitive emissions from solid fuels, Coal mining and handling, pg 8, Table 3-1 have been use for emissions calculating.

NMVOC emission factor is country specific emission factor based on an assessment of the emission factor for methane.

Estimates of emission factors for methane for individual coalmines in Slovenia were done at the Ecological Research Institute (Zapušek A., Orešnik K., Avberšek F: Assessment of methane emission factors in coal excavation in 1986 and in the period 1990-1996, Velenje: ERICO - Ecological Research Institute, 1999). Due to rather small emissions from this sector, no special research project has been done, thus, since 1997, the emission factor recommended in the study period has been assumed. More information on study is presented in Slovenia's National Inventory Report 2016, pg 110.

**Table 3.5.1.2 Emission factors of fugitive emissions in coal mining and handling**

Pollutant	Value	Unit
PM <sub>2.5</sub>	5	g/t
PM <sub>10</sub>	42	g/t
TSP	89	g/t

There is no information whether emission factors for TSP, PM<sub>10</sub>, PM<sub>2.5</sub> represent filterable PM emissions or total (filterable and condensable) emissions.

### Recalculations

No recalculation have been performed since last submission.

### Category-Specific QA/QC and Verification

According to general 2017 in-depth EU NECD review recommendations new EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for emissions calculation. Methodology stated in new guidebook was checked. Since that source is not a key source, Tier 1 method was used for particulate emission calculation. According the EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 the relevant activity statistic for Tier 1 is the total mass of coal produced by underground mining and/or the total tonnage of coal produced by opencast mining.

We consider this approach as an appropriate method for particulate emissions calculation. Since Tier 1 methods in general provide higher emission estimations compared to higher Tier methods, we consider that reported national emissions are therefore not underestimated and completeness of the inventory is assured.

### Future Improvements

No improvement is planned for next submission.

### 3.5.2 Fugitive emissions: Exploration, production and transport of oil and natural gas

NFR Codes covered in this sector:

1B2ai	Fugitive emissions oil: Exploration, production, transport
1B2b	Fugitive emissions from natural gas (exploration, production, processing, transmission, storage, distribution and other)

### Introduction

This chapter deals with the fugitive emissions from the exploration, treatment, loading and also distribution of liquid and gaseous fossil fuels. Oil and natural gas are produced by the same geological process: anaerobic decay of organic matter deep under the Earth's surface. As a consequence, oil and natural gas are often found together. In common usage, deposits rich in oil are known as oil fields, and deposits rich in natural gas are called natural gas fields. Oil and gas are found both onshore and offshore and can be used in a variety of processes, including heating of buildings, and in processes such as feedstock in chemical processes. Natural gas is increasingly being used as a fuel for power generation. The extraction and first treatment of liquid and gaseous fuels involves a number of activities, each of which represents a potential source of hydrocarbon emissions.

Emissions of NMVOC from these sources are insignificant. In 2017 only fugitive emissions from natural gas occurred and contributed less than 0,001 % to total national NMVOC emissions.

### Methodology

To estimate fugitive emissions from production, transport and exploration of oil and natural gas the following methodology has been adopted:

$$E = m \times EF \quad (\text{for crude oil})$$

E – emission (kg)

m – quantity of oil produced (t)

EF – emission factor per quantity of fuel (kg/t)

$$E = m \times EF \quad (\text{for natural gas})$$

E – emission (g)

m – quantity of gas produced (m<sup>3</sup>)

EF – emission factor per quantity of fuel (g/m<sup>3</sup>)

### Activity data

Data on amount of crude oil and natural gas produced have been obtained from SORS. Data for crude oil are given in tonnes. Data for crude oil production is available until 2002. After 2002 there was no production of crude oil. Data on natural gas production are available in the standard m<sup>3</sup> and they are available for the whole 1990-2017 period.

### Emission factors

In calculating emissions of NMVOC emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 3.5.2.1 Emission factors of fugitive emissions**

Pollutant	Value	Unit	Reference
NMVOC (crude oil)	0,2	kg/t	EMEP/EEA Emission Inventory Guidebook, 2016, Fugitive emissions, 1.B.2.a.i Exploration, production, transport, Table 3-1, pg 12
NMVOC (natural gas)	0,1	g/m <sup>3</sup>	EMEP/EEA Emission Inventory Guidebook, 2016, Fugitive emissions, 1.B.2.b Natural gas Table 3-2, pg 12

### Recalculations

No recalculations were performed since last submission.

### Category-Specific QA/QC and Verification

According to general 2017 in-depth EU NECD review recommendations new EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for NMVOC emissions calculation. Methodology stated in new guidebook was checked. Since that source is not a key source, Tier 1 method was used for emission calculation. We consider this approach as an appropriate method for emission calculation. During the review we provided a comparison of current estimations with the estimates resulting with NMVOC emission factors from 2006 IPCC Guidelines. The difference between reported NMVOC emissions and emissions estimated with IPCC EF was insignificant. The impact was far below the threshold of significance. We consider that reported national emissions are therefore not underestimated and completeness of the inventory is assured. We will follow TERT recommendation when EMEP/EEA Guidebook provides emission factors for all segments of natural gas system.

### Future Improvements

No improvement is planned for next submissions.

### 3.5.3 Fugitive emissions oil: Refining / storage

NFR Code 1B2aiv

#### Introduction

This chapter treats emissions from the petroleum refining industry. This industry converts crude oil into more than 2500 refined products, including liquid fuels (from motor gasoline to residual oil), by-product fuels and feedstock (such as asphalt, lubricants, gases, coke), and primary petrochemicals (for instance, ethylene, toluene, xylene). Petroleum refinery activities start with the receipt of crude for storage at the refinery, include all petroleum handling and refining operations, and terminate with storage preparatory to shipping the refined products from the refinery.

Emissions from this source were relevant in Slovenia for 1980-2001 only. Emissions were insignificant and contributed less than 0,0001 % to total national emissions. No emissions of NO<sub>x</sub>, CO, SO<sub>x</sub>, NMVOC, NH<sub>3</sub>, dioxins/furans, heavy metals, particulate matter originated from this sector since 2001.

#### Methodology

To estimate fugitive emissions from refining and storage of oil the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of oil refined (t)

EF – emission factor per quantity of fuel (kg/t)

#### Activity data

Data on amount of crude oil refined have been obtained from SORS. Data for crude oil refined is available until 2001. There was only one oil refinery in Slovenia which was closed down in 2001.

#### Emission factors

In calculating emissions emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 1.B.2.a.iv Fugitive emissions oil: Refining / storage, Table 3-1, pg 14 have been used.

**Table 3.5.3.1 Emission factors of fugitive emissions from refining and storage**

Pollutant	Value	Unit
NO <sub>x</sub>	0,24	kg/t
CO	0,09	kg/t
NMVOC	0,20	kg/t
SO <sub>x</sub>	0,62	kg/t
NH <sub>3</sub>	0,0011	kg/t
PM <sub>10</sub>	0,0099	kg/t
PM <sub>2.5</sub>	0,0043	kg/t
TSP	0,016	kg/t
Cd	0,0051	g/t

<b>Pb</b>	0,0051	g/t
<b>Hg</b>	0,0051	g/t
<b>As</b>	0,0051	g/t
<b>Cr</b>	0,0051	g/t
<b>Cu</b>	0,0051	g/t
<b>Ni</b>	0,0051	g/t
<b>Se</b>	0,0051	g/t
<b>Zn</b>	0,0051	g/t
<b>Dioxins/Furans</b>	0,0057	microg/t

There is no information whether emission factors for TSP, PM<sub>10</sub>, PM<sub>2.5</sub> represent filterable PM emissions or total (filterable and condensable) emissions.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2001.

### Future Improvements

No improvement is planned for next submissions.

## 3.5.4 Distribution of oil products

NFR Code 1B2av

This chapter includes the fugitive emissions of gasoline originating from fuel distribution system. It includes storage in dispatch stations and depots, loading into tank trucks and delivery to the service stations.

### Methodology

To estimate fugitive emissions from distribution of gasoline Tier 2 methodology from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Fugitive emissions, 1.b.2.a.v Distribution of oil products, pg 13 was applied.

### Activity data

Data on amount of gasoline manipulated is obtained from SORS.

### Emission factors

In calculating emissions of NMVOC emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, Fugitive emissions, 1.b.2.a.v Distribution of oil products, Tables 3-2 to 3-12, pg 13-22 have been used.

**Table 3.5.4.1 Emission factors of fugitive emissions in distribution of gasoline**

Pollutant	Value	Unit	Technology	References
NMVOC	23	g/m <sup>3</sup> throughput/kPa TVP	Road tanker	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-4, pg 15
NMVOC	11	g/m <sup>3</sup> throughput/kPa TVP	Rail tanker	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-5, pg 15
NMVOC	24	g/m <sup>3</sup> throughput/kPa TVP	Storage tank filling	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-8, pg 17
NMVOC	3	g/m <sup>3</sup> throughput/kPa TVP	Storage tank breathing	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-9, pg 17
NMVOC	37	g/m <sup>3</sup> throughput/kPa TVP	Automobile refuelling	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-10, pg 18
NMVOC	2	g/m <sup>3</sup> throughput/kPa TVP	Automobile refuelling, drips and minor spilling	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-11, pg 18
NMVOC	0,06	kg/t	Gasoline storage tanks	Emission Inventory Guidebook, 2016, Fugitive emissions, Distribution of oil products, Table 3-12, pg 19

Slovenia implemented Stage I control technique in 2005. Stage II control technique in the refuelling phase was partly implemented in 2010. 51 % of service stations were equipped and operate with Stage II requirements in 2010. In the year 2013 60% of service stations had emission controls for automotive refuelling. Share of service stations with Stage II in 2017 is about 80 %. Abatement efficiencies for vapour recovery were applied for emissions calculation in 2017. For loading facilities this is 98 %, for service stations 95 % and for Stage II automotive refuelling controls 85 %.

### Recalculations

No recalculations have been performed since last submission.

### Category-Specific QA/QC and Verification

According to 2017 in-depth EU NECD review recommendation Tier 2 methodology was applied for emission estimation. Implementation of the control techniques (Stage I and Stage II) was examined and used for emission calculations. New EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for NMVOC emissions calculation.

### Future Improvements

No improvement is planned for next submissions.

### 3.5.5 Venting and flaring (oil, gas, combined oil and gas)

NFR Code 1B2c

#### Introduction

This chapter treats emissions from venting and flaring in the extraction and refining of oil and gas. Flaring is basically combustion of gas, but without utilisation of the energy that is released. Included are flaring during extraction and first treatment of both gaseous and liquid fossil fuels and flaring in oil refineries.

Emissions from this source are insignificant and contributed less than 0,01 % to total national emissions.

#### Methodology

To estimate fugitive emissions from venting and flaring the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – quantity of fuel (t)

EF – emission factor per quantity of fuel (kg/t)

#### Activity data

Data on natural gas produced have been obtained from SORS. Amount of gas burned is 1 % of gas produced.

#### Emission factors

In calculating emissions emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 1.B.2.c Venting and flaring, Flaring in oil and gas extraction, Table 3-1, pg 9 have been used.

**Table 3.5.5.1 Emission factors of fugitive emissions from venting and flaring**

Pollutant	Value	Unit
NO <sub>x</sub>	1,4	kg/t gas burned
CO	6,3	kg/t gas burned
NM VOC	1,8	kg/t gas burned
SO <sub>x</sub>	0,013	kg/t gas burned
PM <sub>10</sub>	2,6	kg/t
PM <sub>2.5</sub>	2,6	kg/t
TSP	2,6	kg/t
BC	0,624	kg/t
Cd	20	mg/t
Pb	4,9	mg/t
Hg	4,7	mg/t
As	3,8	mg/t
Cr	1,3	mg/t



<b>Cu</b>	1,6	mg/t
<b>Ni</b>	38	mg/t
<b>Se</b>	0,43	mg/t
<b>Zn</b>	520	mg/t

There is no information whether emission factors for TSP, PM<sub>10</sub>, PM<sub>2.5</sub> represent filterable PM emissions or total (filterable and condensable) emissions.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-Specific QA/QC and Verification

According to 2017 in-depth EU NECD review recommendation proper activity data were used for NO<sub>x</sub>, CO, SO<sub>x</sub> and NMVOC emission calculation. Emission factors for these pollutants are referred to the gas burned, not to total gas produced. To avoid overestimation we applied new activity data for these pollutants. We use new EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 for emission estimation.

### Future Improvements

No improvement is planned for next submissions.

**Fugitive emission from solid fuels: Solid fuel transformation: NFR Code 1B1b**

**Other fugitive emissions from solid fuels: NFR Code 1B1c**

**Other fugitive emissions from energy production: NFR Code 1B2d**

Notation Key "NO" (not occurring) were used for these three sectors, since there is no other additional fugitive emissions in Slovenia. No emissions occur in these sectors.

## 4 INDUSTRIAL PROCESSES AND PRODUCT USE

Industrial activities not related to energy produce various air emissions. Emission sources are industrial production processes in which raw materials are chemically or physically transformed. In this transformation, many different pollutants into air are released, such as NO<sub>x</sub>, NMVOC, CO, NH<sub>3</sub>, SO<sub>x</sub>, heavy metals and POPs.

Due to the intertwined nature of procedures in industry and characteristics of individual reported units, it is in certain cases difficult to distinguish if certain emissions originate from the consumption of fuels for energy purposes or from the consumption of raw materials in industrial processes. The main criterion is the purpose for which a raw material or fuel is used.

This chapter also deals with the use of paints within the industrial and domestic sectors. It includes emissions arising from degreasing and dry cleaning. It also covers the emissions from the use of chemical products and other solvent use.

According to revised guidelines for reporting emissions and projections data under the Convention LRTAP all emissions from industrial processes and solvent and product use are considered as a whole and reported in one chapter.

### 4.1 Mineral industry (2. A)

Sectors covered in this chapter are:

NFR Codes:

- 2A1 Cement production
- 2A2 Lime production
- 2A3 Glass production

Mineral industry sector contributes to total national emissions with particulate matter and heavy matter emissions. The most important source of emissions of particulate matter in 2017 was lime production. Glass production is the only source of heavy metals. Emissions of TSP and Pb from mineral industry contributed most to national totals, up to 6 % and 3 % respectively.

#### 4.1.1 Cement Production

NFR Code 2A1

During the manufacturing process natural raw materials are finely ground and then transformed into cement clinker in a kiln system at high temperatures. The clinkers are cooled and ground together with additions into a fine powder known as cement. Cement is a hydraulic binder, i.e. it hardens when mixed with water. Cement is used to bind sand and gravel together in concrete.

The basic raw material for the production of cement is marl, which is a homogeneous mixture of limestone and clay and which originated in past geological periods through sedimentation. As there is no longer enough natural marl for mass production, the cement production mix, which must contain 75-78 % of calcium carbonate (CaCO<sub>3</sub>), is prepared by mixing limestone and clay components: from such with 35 % of CaCO<sub>3</sub> to limestone with more than 95 % of

CaCO<sub>3</sub>. The limestone, which is a source of CaO, normally has an admixture of dolomite, which introduces MgO into the system. Clay components are bearers of SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, and Fe<sub>2</sub>O<sub>3</sub>. Blast furnace slag, silica sand, bauxite, and gypsum are added to the homogenized mix during grinding.

Raw meal powder is fed into the cement kiln through a heat exchange unit. Natural gas, fuel oil, petroleum coke, coal dust, waste oils, and tyres are used as fuels in the clinker calcination process.

The present chapter only considers emissions of particulate matter. It comprises emissions of particulate matter from cement plants, which mainly originate from pre- and after-treatment. Emission factors used include also emissions resulting from handling and processing of the product and raw material.

Particulate emissions from combustion process are included in chapter 1.A.2.f. Stationary combustion in manufacturing industries and construction: Non-metallic minerals. As well emissions of NO<sub>x</sub>, SO<sub>x</sub>, CO, NMVOC and NH<sub>3</sub>, heavy metals and persistent organic pollutants from combustion process are included in energy chapter (1.A.2.f.).

Emissions arising in combustion process are calculated from activity data on fuel combusted in cement production and emission factor for particulate fuel and particulate pollutant.

Methodology applied for combustion emission calculation is provided in IIR 2019, Chapter 3.2. Manufacturing industries and construction (1.A.2), pg 68.

In Slovenia, there have been two cement producers until 2015. In the year 2017 only one cement plant has been in operation.

## Methodology

To estimate emissions from cement production, the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of clinker produced (t)

EF – emission factor (kg/t)

## Activity data

Activity data used for emission calculation are data on the annual production of clinker. Data have been obtained from cement producers for the whole period. In 2017 only one cement plant was in operation.

## Emission factors

Emission factors applied for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP and BC emission calculations were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 2A1 Cement production, Table 3-1, pg. 10.

**Table 4.1.1.1 Emission factors for cement production**

Pollutant	Value	Unit	References
TSP	260	g/t	Emission Inventory Guidebook, 2016, 2A1 Cement production, Table 3-1, pg. 10
PM <sub>2.5</sub>	130	g/t	Emission Inventory Guidebook, 2016, 2A1 Cement production, Table 3-1, pg. 10
PM <sub>10</sub>	234	g/t	Emission Inventory Guidebook, 2016, 2A1 Cement production, Table 3-1, pg. 10
BC	3,9	g/t	Emission Inventory Guidebook, 2016, 2A1 Cement production, Table 3-1, pg. 10

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Emissions of particulate matter have been calculated for the period 2000-2017. Emissions from cement production in 2017 contributed up to 1 % to total national emissions.

### Source specific recalculations

No recalculations have been performed since last submission.

### Category-specific QA/QC and verification

Amount of clinker produced and composition of clinker have been thoroughly examined. All data checked were correct. Activity data on clinker production obtained directly from the producers were cross checked with data obtained from verified ETS reports. We also compared data on cement production and clinker production. Clinker production does not entirely track cement production due to additional clinker imports. Cement has been produced not only from domestically produced clinker but also from imported clinker.

According to 2018 in-depth EU NECD review information on methodology applied for combustion emission calculation is provided. Combustion process are calculated from activity data on fuel combusted in cement production and emission factor for particulate fuel and particulate pollutant. Particulate emissions and NO<sub>x</sub>, SO<sub>x</sub>, CO, NMVOC and NH<sub>3</sub>, heavy metals and persistent organic pollutants emissions from combustion process are included in chapter 1.A.2.f. Stationary combustion in manufacturing industries and construction: Non-metallic minerals.

### Planned improvements

No improvements are planned for next submission.

## 4.1.2 Lime Production

NFR Code 2A2

Lime is the high-temperature product of the calcination of limestone. The production occurs in vertical and rotary kilns fired by coal, oil or natural gas. Calcium limestone contains 97–98 % calcium carbonate on a dry basis. Atmospheric emissions in the lime manufacturing industry include particulate emissions from the mining, handling, crushing, screening and calcining of the limestone and emissions of air pollutants generated during fuel combustion in kilns. Lime is

generated by heating the input raw material, i.e. limestone, to high temperature (900-1200°C).

The present chapter only considers emissions of particulate matter.

Combustion related emissions are provided in chapter 1.A.2.f. Emissions arising in combustion process are calculated from activity data on fuel combusted in lime production and emission factor for particulate fuel and particulate pollutant.

In Slovenia, there have been three lime producers until 2013. One of the lime plants had been closed down in the end of 2012. In the year 2017 only two lime plants have been in operation.

### Methodology

To estimate emissions from lime production, the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of lime produced (t)

EF – emission factor (kg/t)

### Activity data

Activity data used for emission calculation are data on the annual production of lime. Data have been obtained from lime producers for the whole period.

### Emission factors

Emission factors applied for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP and BC emission calculations were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 2A2 Lime production, Table 3-1, pg. 8.

**Table 4.1.2.1 Emission factors for lime production**

Pollutant	Value	Unit	References
TSP	9000	g/t	Emission Inventory Guidebook, 2016, 2A2 Lime production, Table 3-1, pg. 8.
PM <sub>2.5</sub>	700	g/t	Emission Inventory Guidebook, 2016, 2A2 Lime production, Table 3-1, pg. 8.
PM <sub>10</sub>	3500	g/t	Emission Inventory Guidebook, 2016, 2A2 Lime production, Table 3-1, pg. 8.
BC	3,22	g/t	Emission Inventory Guidebook, 2016, 2A2 lime production, Table 3-1, pg. 8.

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Emissions of particulate matter have been calculated for the period 2000-2017. Emissions of TSP from lime production in 2017 contributed up to 5 % to total national TSP emissions.

## Recalculations

No recalculations have been performed since last submission.

## Category-specific QA/QC and verification

Amount of lime produced and composition of lime and raw material have been thoroughly examined. Methodology of emission calculation was checked. There were no mistakes found, all data checked were accurate. Activity data on lime production obtained directly from the producers were cross checked with data obtained from verified ETS reports.

## Planned improvements

No improvements are planned for this source.

### 4.1.3 Glass Production

NFR Code 2A3

The present chapter concerns the process emissions released during the production of particular types of glass (flat and container glass, glass wool and Pb glass). It contains emissions for glass production, including emissions from both melting and non-melting activities.

#### Methodology

To estimate emissions from glass production, the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of glass produced (t)

EF – emission factor (kg/t)

#### Activity data

Activity data used for emission calculation are data on the annual production of glass and Pb glass. Data have been obtained from glass producers for the period 2005-2017. For the period 1990-2004 data were obtained from SORS.

#### Emission factors

Emission factors applied for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC, Pb, Cd and Hg emission calculations were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 2A3 Glass production. Emission factors for flat and container glass were taken from Table 3-1, pg. 14, emission factors for lead glass from Table 3-6, pg. 19.

Table 4.1.3.1 Emission factors for glass production

Pollutant	Value	Unit	References
TSP	300	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
PM <sub>2.5</sub>	240	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
PM <sub>10</sub>	270	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
BC	0,1488	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Pb	1,7	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Cd	0,13	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Hg	0,003	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
As	0,19	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Cr	0,23	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Cu	0,007	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Ni	0,49	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Se	0,8	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.
Zn	0,37	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-1 pg. 14.

Table 4.1.3.2 Emission factors for lead crystal glass production

Pollutant	Value	Unit	References
TSP	10	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-6, pg. 19.
PM <sub>2.5</sub>	8	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-6, pg. 19.
PM <sub>10</sub>	9	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-6, pg. 19.
BC	0,00496	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-6, pg. 19.
Pb	10	g/t	Emission Inventory Guidebook, 2016, 2A3 Glass production, Table 3-6, pg. 19.

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Emissions of particulate matter have been calculated for the period 2000-2017 and heavy metals for 1990-2017. Emissions of Pb, Cd, As, Ni, Se contributed up to 3 % to total national lead emissions in 2017.

### Source specific recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### **Category-specific QA/QC and verification**

Amount of glass produced was examined for the whole period. Methodology and emission factors of emission calculation were checked.

### **Planned improvements**

No improvements are planned for next submission.

### **Quarrying and mining of minerals other than coal: NFR Code 2A5a Other mineral products (please specify in the IIR): NFR Code 2A6**

Notation Key "NO" (not occurring) was used for this sector, since there is no quarrying and mining of minerals other than coal in Slovenia. There is also no other mineral products. No emissions occur in these sectors.

### **Construction and demolition: NFR Code 2A5b**

Notation Key "NE" (not estimated) was used for particulate matter in this sector. Emissions of particulates were not estimated since there is no data available for emissions calculation.

### **Storage, handling and transport of mineral products: NFR Code 2A5c**

Emissions of particulate matter from this sector are included under 2A1 Cement production, 2A2 Lime production, 2A3 Glass production. Notation Key "IE" (included elsewhere) was therefore used for this sector.



## 4.2 Chemical industry (2. B)

Sectors covered in this chapter are:

NFR Codes:

2B2	Nitric acid production
2B5	Calcium carbide production
2B6	Titanium dioxide production
2B10a	Chemical industry: Other

Emissions of SO<sub>x</sub> from chemical industry are significant to total national inventory. They contribute 16 % to total emissions. Emissions of other pollutants are negligible. In 2017 only emissions from Titanium dioxide production and Other chemical industry appeared in Slovenia.

### 4.2.1 Nitric acid production

NFR Code 2B2

Nitric acid production is a large scale process in the chemical industry. The process involves the catalytic oxidation of ammonia by air (oxygen) yielding nitrogen oxide then oxidised into nitrogen dioxide (NO<sub>2</sub>) and absorbed in water. The reaction of NO<sub>2</sub> with water and oxygen forms nitric acid (HNO<sub>3</sub>) with a concentration of generally 50–75 wt.% ('weak acid'). For the production of highly concentrated nitric acid (98 wt.%), first nitrogen dioxide is produced as described above. It is then absorbed in highly concentrated acid, distilled, condensed and finally converted into highly concentrated nitric acid at high pressure by adding a mixture of water and pure oxygen.

#### Methodology

To estimate emissions from glass production, the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of nitric acid produced (t)

EF – emission factor (kg/t)

#### Activity data

Activity data for emission calculations are annual production of nitric acid. Data were obtained from Statistical Office of Republic of Slovenia (SORS). Emissions of NO<sub>x</sub> were estimated for the period 1997 – 2005. There is no nitric acid production since 2006.

#### Emission factors

For calculating air emissions from nitric acid production EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 had been used.

**4.2.1.1 Emission factor used for calculation of emissions from nitric acid production**

Pollutant	Value	Unit	References
NO <sub>x</sub>	7,5	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Nitric acid production, Table 3-11, pg. 20

**Emissions**

Since there is no nitric acid production since 2006, no emissions of NO<sub>x</sub> occurred in 2017 from this sector.

**Source specific recalculations**

No recalculation have been performed since last submission.

**Planned improvements**

No improvements are planned for next submission.

**4.2.2 Carbide production**

NFR Code 2B5

Calcium carbide (CaC<sub>2</sub>) is manufactured by heating a lime and carbon mixture up to 2100 °C in an electric arc furnace. The lime is reduced by carbon to calcium carbide and carbon monoxide. Lime for the reaction is usually made by calcining limestone in a kiln at the plant site. The sources of carbon for the reaction are petroleum coke, metallurgical coke and anthracite coal.

**Methodology**

To estimate emissions from calcium carbide production the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of calcium carbide produced (t)

EF – emission factor (kg/t)

**Activity data**

Activity data for emission calculations are annual production of calcium carbide. Data were obtained from SORS. Emissions of TSP were estimated for the period 2000 – 2008. There had been only one producer in Slovenia. This factory was closed down in the first quarter of 2008. There are no emissions from that source since 2008.

### Emission factors

For calculating air emissions from calcium carbide production EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 had been used.

**Table 4.2.2.1 Emission factor used for calculation of emissions from calcium carbide production**

Pollutant	Value	Unit	References
TSP	100	g/t	Emission Inventory Guidebook 2016, Chemical industry, Calcium carbide production, Table 3-5, pg. 16

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Since there is calcium carbide production since 2008, no emissions of TSP occurred in 2017 from this sector.

### Source specific recalculations

No recalculations have been performed since last submission.

### Planned improvements

No improvements are planned for next submission.

## 4.2.3 Titanium dioxide production

NFR Code 2B6

Titanium dioxide (TiO<sub>2</sub>) pigments are made from one of two chemical processes: the chloride route, which leads to TiO<sub>2</sub> products by reacting titanium ores with chlorine gas; and the sulphate route, which leads to TiO<sub>2</sub> products by reacting titanium ores with sulphuric acid. In both processes pure titanium dioxide powder is extracted from its mineral feedstock after which it is milled and treated to produce a range of products designed to be suitable for efficient incorporation into different substrates. This sector represents emissions from sulphate route production in Slovenia.

### Methodology

To estimate emissions from titanium dioxide production the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of titanium dioxide produced (t)

EF – emission factor (kg/t)

### Activity data

Activity data for emission calculations are annual production of titanium dioxide. Data were obtained from SORS until 2016. Data for 2017 have been obtained from the producer.

### Emission factors

For calculating NO<sub>x</sub> and TSP emissions from titanium dioxide production EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used.

**Table 4.2.3.1 Emission factors used for calculation of emissions from titanium dioxide production**

Pollutant	Value	Unit	References
NO <sub>x</sub>	0,108	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Titanium dioxide production, Table 3-20, pg. 25
TSP	0,3	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Titanium dioxide production, Table 3-20, pg. 25

There is no information whether emission factors of particulate matter include or exclude condensable component.

SO<sub>x</sub> emission for the period 2002-2017 are direct emissions taken from REMIS database, established and handled by Slovenian Environmental Agency. These data represent plant specific values.

SO<sub>x</sub> emissions for the period 1982-2001 were estimated. Average EF from 2002-2016 was applied for the period 1982-2001.

For the years 1980 and 1981 emission factor from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used (3,97 kg/t). Abatement technologies started in 1982 and therefore data derived from Remis database was not representative for that period.

The reason for increase in emissions between 2003-2005 is lower efficiency of adsorbent in abatement system.

REMIS database is obtained in compliance with Rules on initial measurements and operational monitoring of the emission of substances into the atmosphere from the stationary pollution sources and on the conditions for their implementation (OJ RS, No. 105/08). Each year all obligators must provide report on implementation of emission monitoring of substances into air. Annual emission report includes emissions of substances into air. These emissions data are direct measurements of emissions into air and reflect plant specific values.

### Emissions

Emissions of SO<sub>x</sub> and NO<sub>x</sub> have been calculated for the period 1980-2017, emissions of TSP for the period 2000-2017. Emissions of SO<sub>x</sub> contributed about 3 % to total national emissions in 2017. Emissions of TSP and NO<sub>x</sub> are below 0,2 %

### Source specific recalculations

According to 2018 in-depth EU NECD review recalculation of SO<sub>x</sub> emissions were performed for the period 1982-2016. Tier 3 plant specific data was used for estimation of SO<sub>x</sub> emissions.

### Category-specific QA/QC and verification

Amount of titanium dioxide produced was examined. Methodology and emission factors of emission calculation were checked. According to 2018 in-depth EU NECD review direct emissions from plants were examined and used for SO<sub>x</sub> emission calculations. Abatement technologies were implemented along 1982-2017 period. Measurements of emissions reflect this improvement, whereas a Tier 2 default emission factor applied for the whole period would not take into consideration such of improvements.

### Planned improvements

No improvements are planned for this source.

## 4.2.4 Chemical industry: Other

NFR Code 2B10a

This sector comprises emissions from formaldehyde, sulphuric acid, polyethylene and NPK (nitrogen, phosphorus, and potassium) and phosphate fertilisers production.

### Methodology

To estimate emissions from other chemical industry production the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of formaldehyde, sulphuric acid, polyethylene or phosphate and NPK fertilisers produced (t)

EF – emission factor (kg/t)

### Activity data

Activity data for emission calculations are annual production of formaldehyde, sulphuric acid, polyethylene and phosphate and NPK fertilisers. Data were obtained from SORS until 2016. Data for 2017 have been obtained from the producer.

### Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emission calculations.

**Table 4.2.4.1 Emission factors used for emissions calculation from formaldehyde production**

Pollutant	Value	Unit	References
NMVOG	1,5	kg/t	Emission Inventory Guidebook, 2016, 2B Chemical industry, Formaldehyde production, Table 3.54, pg. 47

<b>CO</b>	0,2	kg/t	Emission Inventory Guidebook, 2016, 2B Chemical industry, Formaldehyde production, Table 3.54, pg. 47
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**Table 4.2.4.2 Emission factors used for emissions calculation from sulphuric acid production**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>SO<sub>x</sub></b>	3,5	kg/t	Emission Inventory Guidebook, 2016, 2B Chemical industry, Sulphuric acid production, Table 3.25, pg. 27

**Table 4.2.4.3 Emission factors used for emissions calculation from phosphate and NPK fertilizers production**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>TSP</b>	0,3	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Phosphate fertilizers production, Table 3.35, pg. 33
<b>PM<sub>10</sub></b>	0,24	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Phosphate fertilizers production, Table 3.35, pg. 33
<b>PM<sub>2.5</sub></b>	0,18	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Phosphate fertilizers production, Table 3.35, pg. 33

**Table 4.2.4.4 Emission factors used for emissions calculation from polyethylene production**

<b>Pollutant</b>	<b>Value</b>	<b>Unit</b>	<b>References</b>
<b>TSP</b>	0,031	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Polyethylene production, Table 3.39, pg. 37
<b>NM VOC</b>	2,4	kg/t	Emission Inventory Guidebook, 2016, Chemical industry, Polyethylene production, Table 3.39, pg. 37

There is no information whether emission factors of particulate matter include or exclude condensable component.

## Emissions

Emissions of PM<sub>2.5</sub>, PM<sub>10</sub> and TSP from fertilizers and polyethylene production have been calculated for the period 2000 to 2017. Emissions of SO<sub>x</sub> from sulphuric acid production have been calculated for the whole period 1980-2017. Emissions of CO and NMVOC from formaldehyde production had been calculated until 2013. There is no formaldehyde production after year 2014. Sulphuric acid production is significant source of SO<sub>x</sub>. It contributed about 13 % to total national emissions in 2017. Emissions of other pollutants are negligible. They were below 0,1 % of national totals.

## Source specific recalculations

Emissions of SO<sub>x</sub>, NMVOC and TSP have been recalculated for the years 2014 and 2015 due to mistake in calculations made in previous submission.

## Category-specific QA/QC and verification

According to 2018 in-depth EU NECD review this sector was thoroughly examined. No new emissions sources were found. There is a chlorine production in Slovenia, but the process

applied is a membrane cell electrolysis using NaCl. Emissions from this production have not been estimated since no emission factors are available for this type of process. This process has been used in the whole period. Notation key were corrected. 'NE' instead of 'NA' have been used for other pollutants. An error in calculations were found for 2014 and 2015. Mistakes were corrected and recalculations were performed.

### **Planned improvements**

No improvements are planned for next submission.

**Ammonia production: NFR Code 2B1**

**Adipic acid production: NFR Code 2B3**

**Soda ash production: NFR Code 2B7**

**Storage, handling and transport of chemical products: NFR 2B10b**

Notation Key "NO" (not occurring) was used for this sectors, since there is ammonia, adipic acid and soda ash production in Slovenia. No emissions occur in these sectors.

## **4.3 Metal industry (2. C)**

Sectors covered in this chapter are:

NFR Codes:

- 2C1 Iron and steel production
- 2C2 Ferroalloys production
- 2C3 Aluminium production
- 2C5 Lead production
- 2C6 Zinc production
- 2C7a Copper production

The most important source of particulate matter and CO emissions is aluminium production. Steel production is important source of heavy metals and POPs. In 2017 contribution of metal industry to total national emissions is as follows: 33 % to Ni, 25 % to Pb, 24 % to Cd, 23 % to Hg, 15 % to SO<sub>x</sub>, 15 % to dioxins/furans, 11% to Zn and less than 10 % for other pollutants.

### **4.3.1 Iron and Steel Production**

NFR Code 2C1

Iron is produced through the reduction of iron oxide (ore) using metallurgical coke as the reducing agent in a blast furnace. Steel is then subsequently made from iron and scrap in other

furnaces. The production of steel is a multiphase process, and some phases give rise to air emissions. Most emissions occur in smelting iron scrap in electric arc furnace. The furnace is first filled with steel scrap, and then limestone and/or dolomite are added to allow the slag to form. The furnace utilizes electric heating through graphite electrodes. For increased productivity in the initial phase of melting, oxygen lances and a carbon injection system are used. From a metallurgical point of view, oxygen is used to reduce the carbon content in the molten metal and for removing other undesired elements. Decarburising is performed also in secondary phases in a ladle furnace.

There has been only steel production in Slovenia in 2017. Production of pig iron took place until 1987. Since 1988 only electric arc furnace steel plants have been in operation. There have been three steel factories in operation.

Technology of iron and steel production in the period 1980-2017:

1980-1987: pig iron and steel production (integrated plants)

1988-2017: steel production (electric arc furnace)

## Methodology

To estimate emissions from steel production the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of steel produced (t)

EF – emission factor (kg/t)

## Activity data

Activity data used for emission calculation are data on the annual production of steel. For the period 1980-2004 were data obtained from Statistical Office of Republic of Slovenia (SORS). Data on steel produced for the period 2005-2017 have been obtained from steel producers.

## Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emission calculations.

**Table 4.3.1.1 Emission factors used for calculation of emissions from steel production**

Pollutant	Value	Unit	References
TSP	30	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
PM <sub>10</sub>	24	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
PM <sub>2.5</sub>	21	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
BC	0,0756	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
NO <sub>x</sub>	130	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
CO	1,7	kg/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
NM VOC	46	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
SO <sub>x</sub>	60	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron



			and steel production, Table 3.15, pg. 39
<b>Pb</b>	2,6	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Cd</b>	0,2	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Hg</b>	0,05	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>As</b>	0,015	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Cr</b>	0,1	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Cu</b>	0,02	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Ni</b>	0,7	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Zn</b>	3,6Ni,	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>PCB</b>	2,5	mg/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Total 4 PAHs</b>	0,48	g/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39
<b>Dioxins/furans</b>	3	microg I-TEQ/t	Emission Inventory Guidebook, 2016, Metal industry, Iron and steel production, Table 3.15, pg. 39

Emission factors of particulate matter represent filterable emissions only. Condensable component is excluded.

### Emissions

Steel production is important source of heavy metals and POPs. Emissions of Ni, Pb, Cd, Hg contributed about 20 % to national total emissions, emissions of dioxins/furans about 13 %, Zn 11 %, Total 4 PAHs 6 % and PCB 5 %.

### Recalculations

Emissions of As, Cr, Cu, Ni and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Due to change in emission factor used for period 1988-1993 emissions of CO, NO<sub>x</sub> and SO<sub>x</sub> have been recalculated for the period 1988-1993, emissions of NMVOC for the period 1990-1993 and emissions of dioxins/furans, PAHs, PCB, Pb, Cd, Hg for the period 1990-1993.

### Category-specific QA/QC and verification

According to 2018 in-depth EU NECD review recommendation transparency of type of iron and steel technologies occurring in Slovenia was improved. Methodology and emission factors of emission calculation were checked. More appropriate emission factors were used for the period 1988-1993 and recalculations of emissions were performed. Emissions of HCB were not estimated since only electric arc furnace steel plant occurred in the period 1990-2017. No emission factor for HCB is available for this type of process.

### Future improvements

No improvements are planned for next submission.

### 4.3.2 Ferroalloys Production

NFR Code 2C2

Ferroalloys are concentrated alloys of iron and one or more metals such as silicon, manganese, chromium, molybdenum, vanadium and tungsten. These alloys are used for deoxidising and altering the material properties of steel. Ferroalloy production involves a metallurgical reduction process which results in significant carbon dioxide emissions. Emissions from the production of ferroalloys are not considered significant, since the contribution to the total national emissions is thought to be insignificant, i.e. less than 1 % of the national emissions of any pollutant.

#### Methodology

To estimate emissions from ferroalloys production, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of ferroalloys produced (t)

EF – emission factor (kg/t)

#### Activity data

Activity data used for emission calculation are data on the annual production of ferroalloys. Data were obtained from ferroalloys producer for the whole period. This factory was closed down in the first quarter of 2008 and consequently the production of ferroalloys was discontinued in 2008 as well.

#### Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emission calculations.

**Table 4.3.2.1 Emission factors used for calculation of emissions from ferroalloys production**

Pollutant	Value	Unit	References
TSP	1000	g/t	Emission Inventory Guidebook, 2016, Metal industry, Ferroalloys production, Table 3.1, pg. 7
PM <sub>10</sub>	850	g/t	Emission Inventory Guidebook, 2016, Metal industry, Ferroalloys production, Table 3.1, pg. 7
PM <sub>2.5</sub>	600	g/t	Emission Inventory Guidebook, 2016, Metal industry, Ferroalloys production, Table 3.1, pg. 7
BC	60	g/t	Emission Inventory Guidebook, 2016, Metal industry, Ferroalloys production, Table 3.1, pg. 7

Emission factors of particulate matter represent filterable emissions only. Condensable component is excluded.

#### Emissions

Emissions of particulate matter were estimated for the period 2000-2008. There are no emissions from this source since 2008.

## Recalculations

No recalculations have been performed since last submission.

## Future improvements

No improvements are planned for next submission.

### 4.3.3 Aluminium Production

NFR Code 2C3

Aluminium is produced in two phases. Firstly,  $\text{Al}_2\text{O}_3$  is extracted from bauxite ore. Aluminium is then produced in the second phase in an electrochemical process in the electrolysis cells, where alumina disintegrates into its components: aluminium and oxygen. Molten aluminium gathers at the cathode while oxygen reacts with carbon in the anode, causing the consumption of anodes, which have to be replaced. In Slovenia only second phase is performed, when primary aluminium is produced with electrolytic reduction of alumina. In Slovenia, there is one aluminium producer. The most important pollutants emitted from the primary aluminium electrolysis process are sulphur dioxide ( $\text{SO}_2$ ), carbon monoxide (CO), polycyclic aromatic hydrocarbons (PAHs).

## Methodology

To estimate emissions from aluminium production, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of aluminium produced (t)

EF – emission factor (kg/t)

## Activity data

Activity data used for emission calculation are data on the annual production of aluminium. Data have been obtained from aluminium producer for the whole period. Data for primary and secondary aluminium production were used for emission calculation.

## Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for primary and secondary aluminium production emission calculations for:

- $\text{PM}_{2.5}$ ,  $\text{PM}_{10}$ , TSP, BC for the period 2000-2017,
- benzo(a) pyrene, benzo(b) fluoranthene, benzo(k) fluoranthene and Indeno (1,2,3-cd) pyrene, dioxins/furans for the period 1990-2017,
- HCB for the period 1990-2001
- $\text{SO}_x$ ,  $\text{NO}_x$  and CO for the period 1980-1999.

Direct emissions of  $\text{SO}_x$ ,  $\text{NO}_x$  and CO obtained from aluminium producer were applied for the

period 2000-2017.

Since abatement technologies were implemented in secondary aluminium production abatement efficiency of 99 % for dioxins/ furans emission factor in 2017 and 75 % for HCB emission factor in 2001 were used for emissions calculation.

**Table 4.3.3.1 Emission factors used for calculation of emissions from aluminium production**

Pollutant	Value	Unit	References
SO <sub>x</sub>	5	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
NO <sub>x</sub>	1	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
CO	120	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
Benzo(a)pyrene	0,07	g/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
Benzo(b)fluoranthene	0,02	g/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
Benzo(k)fluoranthene	0,02	g/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
Indeno(1,2,3-cd)pyrene	0,01	g/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
TSP	0,6	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
PM <sub>10</sub>	0,5	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
PM <sub>2.5</sub>	0,4	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
BC	0,0092	kg/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.2, pg. 13
Dioxins/ Furans	35	microg I-TEQ/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.4, pg. 15 and Table 3.5, pg. 16
HCB	5	g/t	Emission Inventory Guidebook, 2016, Metal production, Aluminium production, Table 3.4, pg. 15 and Table 3.5, pg. 16

Emission factors of particulate matter represent filterable emissions only. Condensable component is excluded.

According to 2018 in-depth EU NECD review recommendations emissions of HCB and dioxins/furans from secondary aluminium production were introduced into national inventory. In Slovenia there was no use of hexachlorobenzene in secondary aluminium production in the whole period 1990-2017. But there were small amounts of hexachloroethane (HCE tablets) as a degassing agents used for removal unwanted additions in the period 1990-2001. These could cause some small unintentional emissions of HCB in the period 1990-2001. Hexachloroethane was phased out from the production in 2001. HCB emissions were estimated for the period 1990-2001. Emission of dioxins/furans were estimated for the whole period 1990-2017.

## Emissions

Aluminium production is important source of SO<sub>x</sub> and CO. Emissions of SO<sub>x</sub> and CO contributed 13 % and 7 % to total national emissions in 2017. Emissions of other pollutants are less important. They contribute below 0,5 % to national totals. In 2008, a modernisation of technology in aluminium plant was performed. Technological improved point feeding prebaked anode Pechiny has been in operation. A company also acquired the Environmental Permit, which demand introduction of best available techniques and lower the limit of allowed emissions

to the air. For all these reasons, emission factors since 2008 are not comparable with those from years before 2008.

### **Recalculations**

Emissions of dioxins/furans and HCB have been calculated and included into the national inventory for the first time. Emissions of dioxins/furans were introduced for the period 1990-2017, emissions of HCB for the period 1990-2001.

### **Category-specific QA/QC and verification**

According to 2018 in-depth EU NECD review recommendations emissions of HCB and dioxins/furans from secondary aluminium production were introduced into national inventory. Data obtained from aluminium producer was thoroughly examined. Possible inconsistencies were consulted with producer expert team. We also visited the factory and observed production operation and data acquiring in person. Data on direct emissions, which are obtained from producer, are subject to standard QC. In addition implied emission factors are compared with the default EFs from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016. In the cases when IEF is outside the 95% confidence interval we further investigate the reason for such a deviation.

### **Future improvements**

No improvements are planned for next submission.

#### **4.3.4 Lead Production**

NFR Code 2C5

This chapter presents information on atmospheric emissions during primary and secondary lead production. In the direct primary smelting process, the sintering step is skipped, and the lead concentrates and other materials are entered directly into a furnace in which they are melted and oxidized. The secondary production of refined lead amounts to the processing of recycled lead to prepare it for reuse. The vast majority of this recycled lead comes from scrapped lead acid batteries. The most important process emissions are SO<sub>x</sub>, heavy metals and dust.

### **Methodology**

To estimate emissions from lead production, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)  
m – amount of lead produced (t)  
EF – emission factor (kg/t)

### **Activity data**

Activity data used for emission calculation are data on the annual production of lead. Data have

been obtained from SORS until 2016. Data for 2017 was obtained from the producer.

### Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emissions calculation.

**Table 4.3.4.1 Emission factors used for particulate matter emissions calculations from lead production**

Pollutant	Value	Unit	References
TSP	6	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
PM <sub>10</sub>	5	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
PM <sub>2.5</sub>	2,5	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
PCB	2	microg/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
SO <sub>x</sub>	2050	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
Pb	1,8	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
Cd	0,1	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
Hg	0,1	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
As	0,1	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
Zn	0,6	g/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12
Dioxins/furans	4,5	microg I-TEQ/t	Emission Inventory Guidebook, 2016, Metal production, Lead production, Table 3.1, pg. 12

Emission factors of particulate matter represent filterable emissions only. Condensable component is excluded.

### Emissions

Lead production is a minor source of air pollutant emissions. Emissions of all pollutants from lead production contributed less than 2 % to national totals in 2017.

### Recalculations

Emissions of As and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-specific QA/QC and verification

Methodology and emission factors for emission calculation were checked.

### Future improvements

No improvements are planned for next submission.

#### 4.3.5 Zinc Production

NFR Code 2C6

Zinc is produced from various primary and secondary raw materials. The primary processes use sulphidic and oxidic concentrates, while in secondary processes recycled oxidised and metallic products mostly from other metallurgical operations are employed. The most important process emissions are SO<sub>x</sub>, heavy metals and dust.

#### Methodology

To estimate emissions from zinc production, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of zinc produced (t)

EF – emission factor (kg/t)

#### Activity data

Activity data used for emission calculation are data on the annual production of zinc. Data have been obtained from SORS until 2016. Data for 2017 was obtained from the producer.

#### Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emissions calculation.

**Table 4.3.5.1 Emission factors used for particulate matter emissions calculations from lead production**

Pollutant	Value	Unit	References
TSP	15	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
PM <sub>10</sub>	13	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
PM <sub>2.5</sub>	12	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
PCB	2	microg/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
SO <sub>x</sub>	1350	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
Pb	0,2	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
Cd	0,04	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
Hg	0,04	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
As	0,03	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
Zn	5	g/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12
Dioxins/Furans	5	microg I-TEQ/t	Emission Inventory Guidebook, 2016, Metal production, Zinc production, Table 3.1, pg. 12

## Emissions

Zinc production is negligible source of air pollutant emissions. Emissions of all pollutants from zinc production contributed less than 0,05 % to national totals in 2017.

## Recalculations

Emissions of As and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

## Category-specific QA/QC and verification

Methodology and emission factors for emission calculation were checked.

## Future improvements

No improvements are planned for next submission.

### 4.3.6 Copper Production

NFR Code 2C7a

Secondary copper smelter is defined as any plant or factory in which copper-bearing scrap or copper-bearing materials, other than copper-bearing concentrates (ores) derived from a mining operation, is processed by metallurgical or chemical process into refined copper and copper powder (a premium product). The recycling of copper is the most comprehensive among the non-ferrous metals.

## Methodology

To estimate emissions from copper production, the following methodology has been adopted:

$$E = m \times EF$$

E – emission (kg)

m – amount of copper produced (t)

EF – emission factor (kg/t)

## Activity data

Activity data used for emission calculation are data on the annual production of copper. Data have been obtained from SORS until 2016. Data for 2017 was obtained from the producer.

## Emission factors

EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 has been used for emissions calculation.



**Table 4.3.6.1 Emission factors used for particulate matter emissions calculations from copper production**

Pollutant	Value	Unit	References
TSP	320	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
PM <sub>10</sub>	250	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
PM <sub>2.5</sub>	190	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
BC	0,19	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
PCB	0,9	microg/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
SO <sub>x</sub>	3000	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Pb	19	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Cd	11	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Hg	0,023	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
As	4	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Cr	16	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Cu	32	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Ni	14	g/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10
Dioxins/Furans	5	microg I-TEQ/t	Emission Inventory Guidebook, 2016, Metal production, Copper production, Table 3.1, pg. 10

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Copper production is a minor source of air pollutant emissions. Emissions of Cd contributed about 1 %. Emissions of other pollutants are even smaller.

### Recalculations

Emissions of As, Cr, Cu, Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Category-specific QA/QC and verification

Methodology and emission factors for emission calculation were checked.

### Future improvements

No improvements are planned for next submission

**Magnesium production: NFR Code 2C4**

**Nickel production: NFR Code 2C7b**

**Other metal production: NFR Code 2C7c**

Notation Key "NO" (not occurring) was used for these sectors, since there have been no production magnesium, nickel and other metals in Slovenia. No emissions occur in these sectors.

**Storage, handling and transport of metal products: NFR Code 2C7d**

Emissions of this sector are included under 2C1 Iron and steel production, 2C2 Ferroalloys production, 2C3 Aluminium production, 2C5 Lead production, 2C6 Zinc production, 2C7a Copper production. Notation Key "IE" (included elsewhere) was therefore used for this sector.

## **4.4 Solvents and product use (2.D.3 – 2.G)**

### **4.4.1 Description of source category**

This chapter describes the methodology used for calculating air emissions from solvent and product use in Slovenia. The use of solvents and product, containing solvents results in emissions of non-methane volatile organic compounds (NMVOC) when emitted into the atmosphere. In addition to NMVOC emissions, this sector also includes the emissions of other air pollutants as presented in the Table 4.4.1.1.

The most common method of estimating NMVOC emissions is the use of emissions factors. The emissions are estimated based on the production or activity level of the source from which an emission level is calculated using existing Tier 1 or Tier 2 emission factors. The main database of emission factors is the EMEP/EEA Air Pollutant Emission Inventory Guidebook 2016 (GB 2016).

According to this guidebook, emissions from the solvents and other product use are divided into ten sub-categories:

- domestic solvent use including fungicides (NFR 2D3a),
- road paving with asphalt (NFR 2D3b),
- asphalt roofing (NFR 2D3c),
- coating application (NFR 2D3d),
- degreasing (NFR 2D3e),
- dry-cleaning (NFR 2D3f),
- chemical products (NFR 2D3g),
- printing (NFR 2D3h),
- other solvent use (NFR 2D3i), and
- other product use (NFR 2G).

**Table 4.4.1.1: Air pollutants and methodology used for calculation emissions from solvents and other product use in 2017**

NFR	Description	Pollutants	Methods
2D3a	Domestic solvent use including fungicides	NMVOC, Hg	Tier 1
2D3b	Road paving with asphalt	NMVOC, PM	Tier 1 Tier 3
2D3c	Asphalt roofing	NMVOC, CO	Tier 3
2D3d	Coating applications	NMVOC	Tier 3
2D3e	Degreasing	NMVOC	Tier 3
2D3f	Dry cleaning	NMVOC	Tier 3
2D3g	Chemical products	NMVOC	Tier 1, Tier 3
2D3h	Printing	NMVOC	Tier 3
2D3i	Other solvent use	NMVOC, PM, PAHs	Tier 1, Tier 3 Tier 3 Tier 1
2G	Other product use	NMVOC, NO <sub>x</sub> , SO <sub>x</sub> , NH <sub>3</sub> , PM, CO, Pb, Cd, Hg, As, Cr, Cu, Ni, Zn, PCDD/F, PAHs	All pollutants are calculated with Tier 1

In 2017, the solvent and other product use category was the largest source of NMVOC emissions, accounted for 27.3 % of the total NMVOC emissions in Slovenia. The main source is coating application (32.3 %), following by domestic solvent use (30.5 %) and chemical products (27.9 %) while all other sub-categories have contributed only 9.3 % of NMVOC emissions.

Since 1990, NMVOC emissions have decreased by 42.6 % (Figure 4.4.1.1, Table 4.4.1.2) and the largest contribution to this decrease has the decrease of NMVOC emissions from coating application by 64.4%. Two important factors, which have influencing the trend of NMVOC, are the economic situation and environmental legislation.

**Table 4.4.1.2: NMVOC emissions in kt in the period 1990-2017 and relative change of emissions in 2017 to emissions in 1990 and 2016**

	1990	1995	2000	2005	2010	2015	2016	2017	Change to 1990	Change to 2016
2D3a	2.398	2.385	2.388	2.401	2.459	2.476	2.477	2.479	3.4%	0.1%
2D3b	0.012	0.019	0.028	0.024	0.029	0.025	0.025	0.029	133.4%	13.3%
2D3c	0.001	0.001	0.003	0.003	0.001	0.001	0.000	0.000	-72.0%	-28.1%
2D3d	7.385	4.160	5.832	5.440	3.793	2.382	2.402	2.632	-64.4%	9.6%
2D3e	0.203	0.203	0.203	0.203	0.060	0.020	0.033	0.023	-88.4%	-29.8%
2D3f	0.029	0.029	0.029	0.029	0.017	0.007	0.006	0.007	-75.1%	14.1%
2D3g	2.635	2.768	3.684	4.204	3.573	2.122	2.207	2.267	-13.9%	2.8%
2D3h	0.900	0.900	0.900	0.910	0.635	0.200	0.205	0.221	-75.4%	7.8%
2D3i	0.375	0.319	0.344	0.308	0.255	0.197	0.220	0.222	-40.9%	0.7%
2G	0.241	0.242	0.268	0.271	0.261	0.237	0.243	0.255	5.9%	5.0%
<b>Total</b>	<b>14.179</b>	<b>11.027</b>	<b>13.679</b>	<b>13.793</b>	<b>11.083</b>	<b>7.667</b>	<b>7.820</b>	<b>8.137</b>	<b>-42.6%</b>	<b>4.1%</b>

In the period 1990-1993, a reduction of emissions was recorded due to the economic conditions at that time. Slovenian economy went through a variety of shocks in the late 1990s caused by

the transformation of political and economic systems. The crisis was intensified by the loss of former Yugoslav markets. All this resulted in a fall in GDP, a fall in the employment rate and investments, and a high inflation rate. As early as 1993, the Slovenian economy began to revive and the successful economic development lasted to the late 2008 when global financial and economic crisis influenced the first decrease of GDP after 2<sup>nd</sup> quarter of 1993. In the last few years, the economic situation is improving again.

In the May 2004 Slovenia became a member of EU and for this reason have to implement all relevant EU environmental legislation. In the same year, the EU complemented the set of measures to reduce volatile organic matter emissions through Directive 2004/42 / EC on the limitation of emissions of volatile organic compounds due to the use of organic solvents in certain paints and varnishes and vehicle refinishing products. The directive limits the maximum permissible content of volatile organic substances in certain paints and varnishes. Slovenia has implemented this directive with two decrees:

- Decree on limit values for atmospheric emissions of volatile organic compounds from installations using organic solvents (OJ RS, No. 112/05, 37/07, 88/09, 92/10, 51/11, 35/15) and
- Decree on the emission limit values of halogenated volatile organic compounds into the atmosphere from installations using organic solvents (OJ RS, No. 71/11).

According to the VOC legislation every year all VOC obligators must prepare a solvent balance for previous year, taking into account the input and output of solvents, not only through captured and fugitive emissions, but also the proportion of solvents in products and waste. Limit emission values are set for both captured and fugitive emissions of volatile organic substances. The operators from different activities may fulfil their obligations by collecting and purifying volatile organic substances or by implementing an approved plan to reduce emissions of volatile organic substances. Emission reduction plans for volatile organic substances usually involve the transition to the use of paints and varnishes containing a small proportion of volatile substances, as well as more careful solvent management. Since 2005, all data from solvent balance are available in **HOS (VOC) database** and used for estimation of NMVOC emissions from solvent use. Administrator of this database is Slovenian Environmental Agency (SEA).

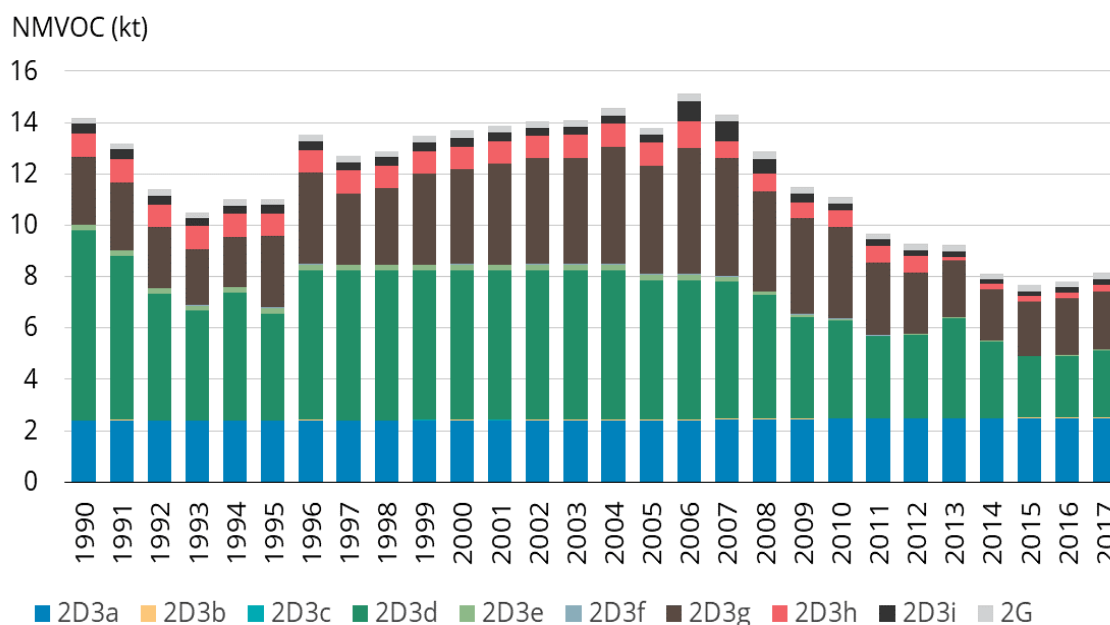


Figure 4.4.1.1: NMVOC emissions from different NFR sub-categories in kt in the period 1990-2017

Besides HOS database the important database that is also located at SEA is a **REMIS database**. Data in the REMIS database are obtained in compliance with Rules on initial measurements and operational monitoring of the emission of substances into the atmosphere from the stationary pollution sources and on the conditions for their implementation (OJ RS, No. 105/08). Each year all obligators must provide report on implementation of emission monitoring of substances into air. These emissions data are direct measurements of emissions into air and reflect plant specific emissions values. In this chapter majority of PMs emissions have been taken from this source and are classified as filterable particulates. Please note that the filterable particulate also includes any material that condenses at or above the filtration temperature.

Due to the large contribution of NMVOC emissions from solvent use to total NMVOC emissions in Slovenia, the peer review of this category has been performed in the late 2016. The results of the peer review and relevant recommendations from the NECD review in 2018 have been taken into account to the extent possible and many improvements have been done for this submission. However, there are still some improvements needed, which are more time demanding and thus are planned for the future submissions. The methodology used and descriptions of recalculations are included in the chapters below under the relevant sub-category.

#### 4.4.2 Domestic solvent use including fungicides

NFR Code 2D3a

This chapter addresses non-methane volatile organic compound (NMVOC) emissions from the use of solvent-containing products by inhabitants in their homes. NMVOCs are used in a large number of products sold for use by the public:

- cosmetics and toiletries,
- household products,
- construction/DIY,
- car care products.

This category does not include the use of decorative paints, which is covered under 2D3d Coating application.

#### Methodology

To estimate emissions from domestic solvent use, the Tier 1 methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the domestic solvent use

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

#### Activity data

Activity data was obtained from Statistical Office of Republic of Slovenia (SORS). In this case, activity data is a number of inhabitants in the Republic of Slovenia on the 1<sup>st</sup> July in particulate year.

## Emission factors

Emissions have been calculated using Tier 1 emission factors from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in the Table 4.4.2.1.

**Table 4.4.2.1 Emission factors used for calculation of NMVOC and Hg emissions from domestic solvent use**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.a
NMVOC	1.2	kg/capita/year	Table 3-1, pg. 8. (other EU countries)
Hg	5.6	kg/capita/year	Table 3-1, pg. 8.

## Recalculations

No recalculations have been performed since the last submission.

## Future improvements

To calculate NMVOC emissions from domestic use of solvents with Tier 2 approach at least the following data on product or solvents use is needed:

- Cosmetics and toiletries: hair sprays, toilet waters, after shaves, perfumes, face cares, personal deodorants and antiperspirants, body care products
- Household products: polishes and creams, soaps
- Car care products: antifreeze

National data on the use of these products is not available.

In the 2016 guidelines, it is proposed that the Tier 2 method has to be used in the EU countries for the assessment, as there are data on the use of solvents in the solvent manufacturer database. In the document (ESIG, 2015), which is indicated in the instructions as a source for data on the use of solvents of solvent-containing products, Slovenia and Austria are taken into account together, therefore data for Slovenia is not available. Because GDP per capita in Slovenia is much smaller than in Austria (in 2017 23,596\$ and 47,221\$ per capita, respectively), comparison with Austria is not suitable.

In addition, the ESIG inventory suggests an emissions factor of 1.2 kg/capita to be used for the EU-27. This EF is used in our calculation of NMVOC emissions.

We will try to estimate activity data, using per capita data from similar country (or group of countries) for the next submission.

### 4.4.3 Road paving with asphalt

NFR Code 2D3b

Asphalt is commonly referred to as bitumen, asphalt cement, asphalt concrete or road oil and is mainly produced in petroleum refineries. Asphalt roads are a compacted mixture of aggregate and an asphalt binder. Natural gravel, manufactured stone (from quarries) or by-products from metal ore refining are used as aggregates. Asphalt cement or liquefied asphalt may be used as the asphalt binder.

## Methodology

To estimate emissions from process of road paving with asphalt, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the road paving with asphalt

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

## Activity data

Since 1998, data on asphalt production is available from the Slovenian Asphalt Pavement Association (<http://www.zdruzenje-zas.si/>), while for the years before, SORS data have been used. In the past data from both sources were similar, but in recent years asphalt production from SORS are distinctively lower from production as reported by association, however the later data looks much more reliable.

## Emission factors

NM VOC emissions have been calculated using Tier 1 emission factors from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.3.1. Emissions of PMs for the period 2000-2004 emissions have been calculated using lower value of Tier 1 emission factor from EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.3.1. Since 2005 measurements of TSP from asphalt plants are available in the Remis database.

**Table 4.4.3.1 Emission factors used for calculation of NM VOC and PM emissions from road paving with asphalt**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.b	Condensable component
NM VOC	16	g/t	Table 3-1, pg. 8	
TSP	10	g/t	Table 3-1, pg. 8. – lower value	excluded
PM <sub>10</sub>	4	g/t	Table 3-1, pg. 8. – lower value	excluded
PM <sub>2.5</sub>	1	g/t	Table 3-1, pg. 8. – lower value	excluded
BC	0.028	g/t	Table 3-1, pg. 8. – lower value	

TSP implied EF for 2005 was 8.8 g/t what is comparable with 10 g/t what is used for the years before. Due to the increasing environmental standards, TSP emissions are decreasing and IEF in 2017 was 2.6 g/t.

As only TSP emissions are available from measurements, other PM emissions have been calculated with the same ratio with TSP as for the years before 2005:

$$E_{\text{PM}_{10}} = 0.4 * E_{\text{TSP}}, \quad E_{\text{PM}_{2.5}} = 0.1 * E_{\text{TSP}}, \quad \text{and} \quad E_{\text{BC}} = 0.0028 * E_{\text{TSP}}$$

Emissions of NO<sub>x</sub>, SO<sub>x</sub>, and CO are expected to originate mainly from combustion and are therefore reported in the category 1.A.2.g.

## Recalculations

No recalculations have been performed since the last submission.

## Future improvements

No improvement is planned for this category.

### 4.4.4 Asphalt roofing

NFR Code 2D3c

Asphalt felt, roofing and shingle manufacture involves the saturation or coating of felt. Heated saturant and/or coating asphalt is applied through dipping and/or spraying. Key steps in the process include asphalt storage, asphalt blowing, felt saturation, coating and mineral surfacing.

#### Methodology

To estimate emissions from Asphalt roofing process, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the asphalt roofing

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

#### Activity data

There is only one plant which produces asphalt roofing product in Slovenia. They are producing asphalt felts and production process is as follows. First the bitumen mass is prepared, depending on the specific product. This bitumen mass is mixed in five 10 m<sup>3</sup> bitumen mixers, using screwed mixers. Mixing takes place at a temperature range of 150 °C to 200 °C. After the appropriate homogeneity of the bitumen mass is reached, it is pumped forward into a horizontally deposited bath, in front of which the carrier (mostly fiberglass) is pre-prepared, winding in the bale. The carrier runs through the inflatable tub where the bitumen mass is applied on the carrier on the both sides of the tape. In the end, the felt is dip in the cooling bath filled with water.

Activity data were obtained from SORS. Data was available in m<sup>2</sup> and for further calculation, we have assumed that 1 m<sup>2</sup> of felt weighted 3 kg. Since 2009 NMVOC emissions has been taken from the Remis database. In the same database PM are not reported as they are negligible and therefore notation key NE is used. Since 2017, emissions of CO are calculated from NMVOC emissions using ratio between CO and NMVOC EF (9.5:46).

#### Emission factors

NMVOC, CO, and PM emission factors were obtained from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook, 2016, as presented in the Table 4.4.4.1. These Tier 2 emission factors are suitable for the production process that is supposed to be in Slovenia: dip saturator, drying in drums section, wet looper and coater.

In the 2016 GB two options of the Tier 2 EFs are available. Tier 2 EFs from the table 3.3, which are appropriate for processes where spray is used are identical as Tier 1 EFs. Because in the



production process in Slovenia no spray is used we have chosen Tier 2 EFs from the Table 3.2 as the most appropriate one. These EFs are used until 2008.

**Table 4.4.4.1 Emission factor used for calculation of emissions from asphalt roofing until the year 2008**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.c
NMVOG	46	g/t shingle	Table 3-2, pg. 8
CO	9,5	g/t shingle	Table 3-2, pg. 8
PM <sub>2.5</sub>	30	g/t shingle	Table 3-2, pg. 8
PM <sub>10</sub>	150	g/t shingle	Table 3-2, pg. 8
TSP	600	g/t shingle	Table 3-2, pg. 8
BC	0,0039 (0.013% of PM <sub>2.5</sub> )	g/t shingle	Table 3-2, pg. 8

*Note: There is no information in the EMEP/EEA GB 2016 whether EFs of PM include or exclude condensable component.*

### Recalculations

Since 2009 NMVOG emissions have been recalculated using data from the Remis database.

### Future improvements

No improvement is planned for this category.

## 4.4.5 Coating Application

NFR Code 2D3d

The use of paint is a major source of NMVOG emissions; they comprise almost 10 % of total NMVOG emissions in the country. The use of paints is generally not considered relevant for emissions of particulate matter or heavy metals and POPs. Most paints contain organic solvent, which must be removed by evaporation after the paint has been applied to a surface in order for the paint to dry or 'cure'. Unless captured and either recovered or destroyed, these solvents can be considered to be emitted into the atmosphere. Some organic solvent may be added to coatings before application, which will also be emitted. Further solvent used for cleaning coating equipment is also emitted.

The proportion of organic solvent in paints can vary considerably. Traditional solvent borne paints contain approximately 50 % organic solvents and 50 % solids. In addition, more solvent may be added to further dilute the paint before application. High solids and waterborne paints both contain less organic solvent, typically less than 30 %, while powder coatings and solvent free liquid coatings contain no solvent at all. NMVOG emissions, which are calculated using EF, are thus less accurate than measured emissions, which are also used in this category.

The main source of NMVOG emissions in this category is decorative coating application. It could be applied by enterprises and professional painters (SNAP activity 060103) or by private consumers (SNAP activity 060104). For inventory purpose distinguish between both types of uses was not possible.

In this category the following industrial coating application are also included:

Manufacture of automobiles (SNAP activity 060101)

This category refers to the coating of automobiles as part of their manufacture; it includes corrosion protection at point of manufacture. The application of sealants as part of the manufacturing process is covered here.

Car repairing (SNAP activity 060102)

This category refers to the coating of road vehicles carried out as part of vehicle repair, conservation or decoration outside of manufacturing sites, or any use of refinishing-type coatings where this is carried out as part of an original manufacturing process.

Coil coating (SNAP activity 060105)

This category refers to the coating of coiled steel, aluminium or copper alloy strips as a continuous process.

Boat building (SNAP activity 060106)

This category refers to all paints for the hulls, interiors and superstructures of both new and old ships and boats.

Wood (SNAP activity 060107)

Wood may be colour coated, stained or varnished and the fugitive emissions could be significant.

Other industrial paint application (SNAP activity 060108)

This category refers to all industrially applied paints for metal, plastic, paper, leather and glass substrates, which are not covered by any of the other categories described above.

**Methodology**

To estimate emissions from coating application, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

Since 2005 NMVOC emissions from the industrial sources have been taken from the HOS database and since 2011 the total NMVOC emissions in this category is determined using “tax database”.

**Activity data**

Activity data for NMVOC emission calculations from industrial coating application for the period 1990 to 1996 were obtained from SORS. After the year 1996 SORS did not provide paint consumption data at all. Therefore, the emission values from the year 1996 have been used until the year 2004. Since 2005 NMVOC emissions from the HOS database have been used. In the previous submission, we have used Tier 1 approach and constant factor of 6.7 kg paint/capita/year to estimate amount of paint used for the decorative coating application. This approach has been also recommended in the expert peer review.

For the present submission, we have obtained data on amount of VOC in different products. This amount has been collected for the determination of environmental tax, which is payable for paints and varnishes and products for lacquering motor vehicles (Decree on environmental tax for environmental pollution due to use of volatile organic compounds). Tax payers are producers and acquirers from Slovenia and other EU countries or from third countries, if their annual

quantity of acquired or produced above mentioned products exceeds 150 kg. The tax is payed for amount of VOC in these product and the data are available since 2010. This tax is covering all product regardless of whether they are intended for domestic use or to be used in the industry. For this reason, we have assumed that a yearly amount of VOC in these products is the same as yearly emissions of NMVOC emissions from coating application.

To validate this assumption we have compared NMVOC emissions for 2010 using the old and the new approach and difference was less than 3%. We have chosen the year 2010 because the amount of paint per person in the GAINS model was determined for this year. We believe that the data from the tax database are reliable and we are using them for determination of NMVOC emissions since 2011.

We have also try to use Eurostat data as was recommended in 2017 NECD review but resulted estimate of decorative paint was unreasonable high and the fluctuations between years was even higher.

### Emission factors/ Emissions

Until 2010 NMVOC emissions from the decorative coating applications have been calculated using Tier 1 emission factors from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.5.1.

NMVOC emission factor for industrial coating application in the period 1990 to 1996 were also obtained from EMEP/EEA air pollutant emission inventory guidebook 2016. Emissions of NMVOC from the year 2005 onwards have been taken from HOS database.

**Table 4.4.5.1 Tier 1 emission factor used for calculation of NMVOC emissions from decorative coating application**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.d
NMVOC	150	g/kg paint applied	Table 3-1, pg. 17
NMVOC	400	g/kg paint applied	Table 3-2, pg. 17

A detailed overview of the emission determination for each period can be found in the table 4.4.5.2 below.

**Table 4.4.5.2 Overview of determination of NMVOC emissions and sources of data in different periods**

Coating application	1990-1996	1997-2004	2005-2010	Since 2011
Domestic	6.7 kg paint/cap * EF	6.7 kg paint/cap * EF	6.7 kg paint/cap * EF	Calculated: (Total - Industrial)
Industrial	Paint use (SORS) * EF	Emission from 1996	Emissions from the HOS database	Emissions from the HOS database
Total 2D3d	sum	sum	sum	Amount of VOC from the TAX database

### Source specific recalculations

For the period 2011-2016 NMVOC, emissions have been recalculated using data on amount of NMVOC from the tax database.

### **Source-specific planned improvements**

For the next submission, we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

The next step of improvement for this category would be a split of decorative coating application between domestic use and paint use in construction and buildings. At that moment, we have no data and no reliable methodology to perform such disaggregation. As this improvement would have no effect on the total emissions, it is not planned for the near future.

### **4.4.6 Degreasing**

NFR Code 2D3e

Degreasing is a process for cleaning products from water-insoluble substances such as grease, fats, oils, waxes, carbon deposits, fluxes and tars. In most cases, the process is applied to metal products, but also plastic, fibreglass, printed circuit boards and other products are treated by the same process.

#### **Emission factors / Emissions**

Emissions of NMVOC from the year 2005 onwards have been taken from HOS database. Emissions of NMVOC for the period 1990-2004 were estimated, since no data are available before the year 2005.

#### **Recalculations**

No recalculations were performed since the last submission.

### **Future improvements**

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

### **4.4.7 Dry Cleaning**

NFR Code 2D3f

Dry cleaning can be defined as the use of chlorinated organic solvents, principally tetrachloroethene, to clean clothes and other textiles. In general, the process can be divided into four steps:

- cleaning in a solvent bath,
- drying with hot air and recovery of solvent,
- deodorisation (final drying),
- regeneration of used solvent after the clothes have been cleaned.

### Emission factors/ Emissions

Emissions of NMVOC from the year 2005 onwards have been taken from HOS database. Emissions of NMVOC for the period 1990-2004 were estimated, since no data are available before the year 2005.

### Recalculations

No recalculations were performed since the last submission.

### Future improvements

No improvements are planned for this category.

## 4.4.8 Chemical Products

NFR Code 2D3g

Emission sources of NMVOC in Slovenia are generated during the manufacturing of the following products:

- Polyvinyl chloride and other plastic (SNAP 060301-4)
- Rubber products (SNAP 060305)
- Pharmaceutical products (SNAP 060306)
- Paints (SNAP 060307)
- Inks (SNAP 060308)
- Glues (SNAP 060309)
- Leather tanning (SNAP 060313)

### Methodology

To estimate emissions from chemical products, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

### Activity data

#### Polyvinyl chloride and other plastic

Statistical data distinguish between Plastics in primary forms (C20.16) and Plastics products (C22.2). According to this data majority of primary plastics is further shaped in the plastic products, what means that inclusion of this amount in the calculation will lead to double counting of emissions. For this reason emissions from plastics product (remaking of plastics) have been excluded from the inventory. This recommendation has been also included in the final report of the peer review in 2016 (Česen, 2016).

Data on the production of plastics has been obtained from SORS under special memorandum and with awareness that this are confidential data, because number of producers for every type of product is very limited. Data are available in tonnes for each code from NIP (National

nomenclature of Industrial Products). More information on NIP is available here (<http://www.stat.si/StatWeb/en/Methods/Classifications>). For determination of the NMVOC emissions all types of plastics which are reported under code C20.16 Plastics in primary forms have been sum together and multiply with Tier 1 EF of 10 g NMVOC/kg product.

According to the 2016 EEA/EMEP 2016 only four plastic processing have to be included in the inventory: Polyester- PE, Polyvinylchloride - PVC, Polyurethane foam - PUR and Polystyrene foam - PS. In statistical data only production of PE and PUR is available, while data on PS and PVC are not available. There is also no Tier 2 EF for PVC production in the 2016 EMEP/EEA GB while Tier 2 EF for PE production is available for amount of monomer used and not for PE produced. For all these reasons we didn't use Tier 2 EFs for 2019 submission yet. In the further investigation we have found out that CO<sub>2</sub> has been used as a blowing agent for the PUR production since 1994 in one plant. On the other hand despite data on PS production are not available from the SORS there are three PS foam producers in Slovenia.

The expert in peer review recommended that measurements of NMVOC (total organic o-toluidine) from relevant processes are the best option to estimate emissions from these sources. NMVOC emissions from these plants are available since 2009. In 2016 emissions from these plants were only 0.078 kt, while emissions from production of plastics calculated with AD (Plastics in primary forms) \* Tier 1 EF were 1.45 kt. The emission for other years have not been obtained yet. We will investigate this issue more precisely for the next submission.

#### Rubber products

Data on rubber products for the period 1990-2004 have been obtained from the SORS. Under this category all rubber production is included. Because majority of the rubber products were tyres, the Tier 2 EF for manufacture of tyres has been used, while since 2005 emissions from the HOS database have been used.

#### Pharmaceutical products

Emissions from pharmaceutical products are included in the inventory, data since 2005 has been taken from HOS database.

#### Paints, Inks, and Glues

Data on production were obtained from SORS for all years and NMVOC emissions are calculated using Tier 2 EF.

#### Leather tanning

Emissions from leather tanning are included in the inventory, data since 2005 has been taken from HOS database. Since 2009 there is no more Leather tanning industry in Slovenia. The leather industry of Vrhnika (IUV) was the largest European leather factory. However, in December 2008, the company announced bankruptcy due to a number of unfavorable circumstances linked to global financial crisis and domestic economic and political dynamics.

Emissions from other chemical products are not occurring in the country. There is no asphalt blowing process in Slovenia. We have no oil refinery, and this process is not used in the asphalt processing or asphalt roofing plants in Slovenia. Total amount of the air-blown bitumen which is used in the production of asphalt roofing product is imported.

### **Emission factors/ Emissions**

NMVOC emissions from the production of chemical products have been calculated using Tier 1 and Tier 2 emission factors from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.8.1.

**Table 4.4.8.1 Emission factors used for calculation of NMVOC emissions from chemical products**

	Unit	Value	Source: GB 2016, NFR 2.D.3.g
Plastics	kg/t	10	Table 3-1 (Tier 1)
Rubber products	kg/t	10	Table 3-6 (Tier 2)
Oil paints and inks	kg/t	11	Table 3-11 (Tier 2)
Glue	kg/t	11	Table 3-11 (Tier 2)

Since 2005 emissions of NMVOC from paints and rubber processing have been taken from HOS database.

### Recalculations

Due to the obvious error in AD for 2001, NMVOC emissions for this year have been interpolated.

### Future improvements

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

Following the TERT recommendation, we will obtain data on NMVOC (total organic o-toluidine) emissions from Remis database since 2009 for relevant plastics production processes and assess the implementation of this methodology for the next submission.

## 4.4.9 Printing

NFR Code 2D3h

Printing involves the use of inks, which may contain a proportion of organic solvents. These inks may then be subsequently diluted before use. Different inks have different proportions of organic solvents and require dilution to different extents. Printing can also require the use of cleaning solvents and organic dampeners, Ink solvents, diluents, cleaners and dampeners.

There is a strong decreasing trend of NMVOC emissions from printing with two sharp drops in 2007 and in 2012. The first one is connected to the implementation of VOC directive, while the second one is influenced with the decline in printed media and increasing use of cleaning devices.

### Activity data

Activity data for NMVOC emission calculations from the year 1990 to 1996 were obtained from SORS. After the year 1996 SORS did not provide paint consumption data at all. Therefore, NMVOC emission data from the year 1996 have been used until 2004. For the period 2005-2016 NMVOC emissions from HOS database have been applied.

### **Emission factors/ Emissions**

NMVOC emission factor for the period 1990 to 1996 were obtained from CORINAIR INVENTORY Default Emission Factors Handbook (second edition), 1992, (EF NMVOC, 200 kg/t).

Since 2005, all the factories in industry and private sector, who use paint and varnish or other solvent are obliged to report their emissions annually and Slovenia considers that their data cover more than 97 % of all emissions from printing industries. For this reason, emissions of NMVOC from the year 2005 onwards have been taken from HOS database.

### **Recalculations**

No recalculations have been performed since the last submission.

### **Future improvements**

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

#### **4.4.10 Other solvent and product use**

NFR Codes 2D3i and 2G

Emission sources covered in this chapter can be divided into two sub-categories:

Sources of emissions from 2D3i other solvent use are:

- Mineral wool production (060402),
- Fat, edible and not edible oil extraction (060404),
- Application of glues and adhesives (060405),
- Preservation of wood (060406),

while under 3G emissions from the following product use have been included:

- Use of fireworks (060601),
- Use of tobacco (060602),
- Use of shoes (060603),
- Other (060604) – Use of pesticides. Airplane de-icing.

Emissions from glass wool production (060401) are included in the category 2A3 Glass production. Emissions from the asphalt blowing do not occur in the country.

Emissions of underseal treatment and conservation of vehicles as well as vehicle dewaxing have been not estimated due to the unavailability of activity data. The expert judgement from the peer review is that emissions from this source in Slovenia are negligible.

#### Mineral wool production

To estimate emissions from mineral wool production the following methodology has been adopted:



$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the mineral wool production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

### Activity data

Activity data on annual production of mineral wool, obtained from SORS, are confidential.

### Emission factors/ Emissions

NMVOC emissions from the mineral wool production have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.1.

**Table 4.1.10.1 Emission factor used for calculation of NMVOC emissions from Mineral wool production for NMVOC**

Pollutant	EF	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NMVOC	300	g/t	Table 3-3

### Fat, edible and not edible oil extraction and Application of glues and adhesives

Emissions of NMVOC from Fat, edible and not edible oil extraction and Application of glues and adhesives from the year 2005 onwards have been taken from the HOS database.

In addition, PM emissions from grain handling process in the oil production have been included. Since 2005, emissions of TSP have been taken from the Remis database, while for the period 2000 to 2004 the 2005 value has been used.

Only emissions of TSP are available from measurements. Thus other PM emissions have been calculated with the same ratio with TSP as presented on the Table 3.4 in EMEP/EEA air pollutant emission inventory guidebook 2016, 2D3i, 2G Other solvent and product use:

$$E_{\text{PM}_{10}} = 0.9/1.1 * E_{\text{TSP}}, E_{\text{PM}_{2.5}} = 0.6/1.1 * E_{\text{TSP}}, \text{ and BC emissions are not estimated (NE).}$$

### Preservation of wood

To protect wood against wood decay fungi and insects and also against weathering, wood preservatives that fully penetrate into wood, need to be applied. In practice, wood preservatives are applied only by brushing. There are three main types of preservative: creosote, organic solvent-based (often referred to as 'light organic solvent-based preservatives) and water borne. Creosote is an oil prepared from coal tar distillation. Creosote contains a high proportion of aromatic compounds such as polycyclic aromatic hydrocarbons (PAHs). Levels of benzo(a)pyrene in some types of creosote are restricted in the EU to 500 ppm as well in Slovenia for industrial use (14th amendment to the Marketing and Use Directive — Creosote (96/60/EEC)).

To estimate emissions from preservation of wood the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant  
 $AR_{\text{production}}$  – the activity rate for the production  
 $EF_{\text{pollutant}}$  – the emission factor for this pollutant

### Activity data

Activity data were obtained from impregnation of wood plant (personal communication).

### Emission factors

NMVOC and PAH emissions from the preservation of wood have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016 for creosote preservative type, as presented in Table 4.4.10.2.

**Table 4.4.10.2 Tier 2 emission factors used for calculation of NMVOC and PAH emissions from wood preservation**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NMVOC	105	kg/t	Table 3-5
Benzo(a)pyrene	1.05	g/t	Table 3-5
Benzo(b)fluoranthene	0.53	g/t	Table 3-5
Benzo(k)fluoranthene	0.53	g/t	Table 3-5
Indeno(1,2,3-cd)pyrene	0.53	g/t	Table 3-5

### Use of fireworks

#### Activity data

The quantity of used fireworks in Slovenia (Table 4.4.10.3) is estimated by the import and export data (CN codes 36041000 and 36049000) available from Eurostat Database. There is no production of fireworks in Slovenia. Data regarding import and export are not available for the years 1990-1998 and emissions for this period are estimated to be similar as in 1999.

**Table 4.4.10.3 Activity data for fireworks**

Year	Fireworks (t)	Year	Fireworks (t)
1990-1998	250.0	2011	456.1
1999	243.0	2012	720.7
2000	203.8	2013	307.8
2001	265.8	2014	183.0
2002	317.1	2015	467.8
2003	407.7	2016	442.6
2004	493.8	2017	290.8
2005	926.0		
2006	629.6		
2007	464.2		
2008	773.4		
2009	181.3		
2010	628.8		

## Emission factors

Air pollutant emissions from the use of fireworks have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.4.

**Table 4.4.10.4 Emission factors used for calculating pollutant emissions from the use of fireworks**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
SO <sub>2</sub>	3,020	g/t	Table 3-13
NO <sub>x</sub>	260	g/t	Table 3-13
CO	7,150	g/t	Table 3-13
TSP	109,830	g/t	Table 3-13
PM <sub>10</sub>	99,920	g/t	Table 3-13
PM <sub>2.5</sub>	51,940	g/t	Table 3-13
As	1.33	g/t	Table 3-13
Cd	1.48	g/t	Table 3-13
Cr	15.6	g/t	Table 3-13
Cu	444	g/t	Table 3-13
Hg	0.057	g/t	Table 3-13
Ni	30	g/t	Table 3-13
Pb	784	g/t	Table 3-13
Zn	260	g/t	Table 3-13

Note: There is no information in the EMEP/EEA GB 2016, whether EF of PM include or exclude condensable component.

## Tobacco combustion

### Activity data

The quantity of tobacco combusted in Slovenia (Table 4.4.10.5) has been taken from the WHO study [Tobacco taxation policy in Slovenia](#) and for recent years from the [Tobacco atlas](#).

**Table 4.4.10.5 Use of tobacco in tons**

Year	Tobacco (t)	Year	Tobacco (t)
1990-2001	3,750.0	2011	3,825.0
2002	3,600.0	2012	3,900.0
2003	3,487.5	2013	3,975.0
2004	3,375.0	2014	4,077.4
2005	3,412.5	2015	3770.4
2006	3,675.0	2016	3,462.5
2007	3,450.0	2017	3,465.7
2008	3,750.0		
2009	3,900.0		
2010	3,750.0		

## Emission factors

Air pollutant emissions from tobacco combustion have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.6.

**Table 4.4.10.6 Emission factors used for calculating pollutant emissions from tobacco combustion**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NM VOC	4.84	kg/t tobacco	Table 3-14
NO <sub>x</sub>	1.80	kg/t tobacco	Table 3-14
CO	55.1	kg/t tobacco	Table 3-14
NH <sub>3</sub>	4.15	kg/t tobacco	Table 3-14
TSP	27.0	kg/t tobacco	Table 3-14
PM <sub>10</sub>	27.0	kg/t tobacco	Table 3-14
PM <sub>2.5</sub>	27.0	kg/t tobacco	Table 3-14
BC	0.45	% of PM <sub>1.8</sub>	Table 3-14
PCDD/F	0.1	µg I-TEQ/t tobacco	Table 3-14
Benzo(a)pyrene	0.111	g/t tobacco	Table 3-14
Benzo(b)fluoranthene	0.045	g/t tobacco	Table 3-14
Benzo(k)fluoranthene	0.045	g/t tobacco	Table 3-14
Indeno(1,2,3-cd)pyrene	0.045	g/t tobacco	Table 3-14
Cd	5.4	g/t tobacco	Table 3-14
Ni	2.7	g/t tobacco	Table 3-14
Zn	2.7	g/t tobacco	Table 3-14
Cu	5.4	g/t tobacco	Table 3-14

Note: There is no information in the EMEP/EEA GB 2016 whether EF of PM include or exclude condensable component.

#### Use of shoes

##### **Activity data**

It is not clear from the guidebook what should be used as activity data for use of shoes; is this all pair of shoes, bought in one year or all pairs of shoes used in one year? We decided to use population number as no one can use more as one pair of shoes at a time.

##### **Emission factors**

NM VOC emissions from the use of shoes have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.7.

**Table 4.4.10.7 Emission factors used for calculating NM VOC emissions from the use of shoes**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NM VOC	60	g/pair	Table 3-15

#### Other - use of pesticides

##### **Activity data**

Activity data on pesticides used in the country has been obtained from the SORS and are available on the Table 4.4.10.8.

Table 4.4.10.8 Amount of pesticides used in tons

Year	Pesticides (t)	Year	Pesticides (t)
1990-1997	1,267.143	2011	1,121.873
1998	1,115.851	2012	1,016.069
1999	1,605.972	2013	917.483
2000	1,468.110	2014	1,009.912
2001	1,398.268	2015	1,046.822
2002	1,164.089	2016	1,156.192
2003	1,361.003	2017	1,087,286
2004	1,557.980		
2005	1,313.967		
2006	1,280.980		
2007	1,155.221		
2008	1,218.151		
2009	1,162.873		
2010	1,134.370		

### Emission factors

NMVOC emissions from the use of pesticides have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.9.

Table 4.4.10.9 Emission factors used for calculating NMVOC emissions from the use of pesticides

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NMVOC	69,000	g/t	Table 3-16

### Other –airplane de-icing

#### Activity data

Activity data on the use of de-icing agent since 2009 has been obtained from the Airport Ljubljana while the data for the years before was estimated taking into account number of flights and climate conditions. Since 2009 de-icing agents used were Kilfrost DF Plus (Type I) and Kilfrost ABC-S Plus (Type IV). The amount of de-icing agent used since 1990 is available on the Table 4.4.10.10.

Table 4.4.10.10 Amount of de-icing agent used in tons

Year	De-icing agent (t)	Year	De-icing agent (t)	Year	De-icing agent (t)
1990	65.012	2000	119.860	2010	167.864
1991	35.176	2001	116.200	2011	116.081
1992	35.444	2002	114.284	2012	148.930
1993	51.592	2003	126.948	2013	128.187
1994	63.284	2004	142.008	2014	78.674

<b>1995</b>	71.472	<b>2005</b>	151.068	<b>2015</b>	93.996
<b>1996</b>	72.760	<b>2006</b>	163.964	<b>2016</b>	93.761
<b>1997</b>	81.116	<b>2007</b>	186.068	<b>2017</b>	160.636
<b>1998</b>	102.892	<b>2008</b>	191.704		
<b>1999</b>	108.876	<b>2009</b>	189.704		

### Emission factors

NMVOC emissions from the use of de-icing agent have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.4.10.11.

**Table 4.4.10.11 Emission factors used for calculating NMVOC emissions from the airplane de-icing**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.D.3.i, 2.G
NMVOC	246,000	g/t	Table 3-16

### Recalculations

Following the recommendation from the peer review NMVOC emissions from airplane de-icing have been included in the inventory for the period 1990-2017.

In addition, emissions of six heavy metals (As, Cr, Cu, Ni, Se, and Zn) for the period 1990-2017 have been included in the inventory for the first time.

### Future improvements

For category 2D3i Other Solvent Use and pollutant NMVOC for year 2006 the TERT noted that there was a sharp increase of NMVOC emissions in 2006 by 2.8 times compared to the year 2005. After investigation we have found out, that HOS database was not completed for production of glues for 2005, 2006 and 2008. For this reason, we have rechecked all data in the HOS database for all years and all categories and some other missing data have been found. Now, the database is complete, but was not complete at the time of inventory submission on 15 February. Therefore, the recalculation of emissions from HOS database in the year 2005, 2006, and 2008 will be done for the next submission. This recalculation will also affect emissions for the period 1990-2004, because in some cases when the data for the years before 2005 are not available, emissions in the in 2005 was used to extrapolate emissions back to 1990.

## 4.5 Other industry production (2. H)

Emission sources covered in this chapter are:

- 2H1 Pulp and paper industry
- 2H2 Food and beverages industry

No other relevant industrial production has occurred in Slovenia and notation key NO has been used for category 2H3.

#### 4.5.1 Pulp and paper industry

NFR Code 2H1

Paper is essentially a sheet of cellulose fibres with a number of added constituents to affect the quality of the sheet and its fitness for intended end use. The pulp for papermaking may be produced from virgin fibre by chemical or mechanical means or by the re-pulping of recovered paper. In the pulping process, the raw cellulose-bearing material is broken down into its individual fibres. Wood is the main raw material but straw, hemp, grass, cotton and other cellulose-bearing materials can be used as well. The precise composition of the wood will vary according to the type and species but the most important constituents are cellulose, hemicelluloses and lignin. In Slovenia, there were five pulp and paper plants and some of them were closed for operation in some years.

#### Methodology

To estimate emissions from pulp and paper, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

#### Activity data

Activity data on pulp production until 2005 were obtained from SORS, while since then the measurements have been used.

#### Emission factors

For calculating air emissions from pulp and paper until 2005 we have used Tier 2 EFs from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.5.1.1. These EFs are suitable for the Kraft pulping process, which was abolished in 2006 and since then a pulp is produced with a process called thermo-mechanical pulp production, while for bleaching a sulphite or peroxide have been used. No emission factors are available for this type of production in the EMEP/EEA GB 2016, hence since 2006 NMVOC emissions are taken from the Remis database as a sum of emissions of TOC from five pulp and paper producers, while for other pollutants notation keys NA or NE are used.

**Table 4.5.1.1 Emission factors used for calculation of emissions from pulp and paper 1990-2005**

Pollutant	Value	Unit	Source: GB 2016, NFR 2.H.1
NO <sub>x</sub>	1,0	kg/t	Table 3-2
CO	5,5	kg/t	Table 3-2
NMVOC	2,0	kg/t	Table 3-2
SO <sub>x</sub>	2,0	kg/t	Table 3-2
PM <sub>2.5</sub>	0,6	kg/t	Table 3-2
PM <sub>10</sub>	0,8	kg/t	Table 3-2
TSP	1,0	kg/t	Table 3-2
BC	0,0156	kg/t	Table 3-2

*Note: There is no information in the EMEP/EEA GB 2016 whether EFs of PM include or exclude condensable component.*

## Recalculations

Since 2006 emissions of NMVOC have been recalculated using measurements.

## Future improvements

No improvements are planned for this category.

### 4.5.2 Food and beverages industry

NFR Code 2H2

Food manufacturing may involve the heating of fats and oils and foodstuffs containing them, the baking of cereals, flour and beans, fermentation in the making of bread, the cooking of vegetables and meats, and the drying of residues. These processes may occur in sources varying in size from domestic households to manufacturing plants. When making any alcoholic beverage, sugar is converted into ethanol by yeast. This is fermentation. The sugar comes from fruit, cereals or other vegetables. These materials may need to be processed before fermentation. To make spirits, the fermented liquid is then distilled. Alcoholic beverages, particularly spirits and wine, may be stored for a number of years before consumption. Emissions may occur during any of the four stages, which may be needed in the production of an alcoholic beverage. During preparation of the feedstock, the most important emissions appear to occur during the roasting of cereals and the drying of solid residues. During fermentation, alcohol and other NMVOCs are carried out with the carbon dioxide as it escapes to atmosphere. In some cases, the carbon dioxide may be recovered, reducing the emission of NMVOC as a result.

## Methodology

To estimate emissions from food and drink, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

## Activity data

Activity data for emission calculations were obtained from SORS.

The relevant activity statistics are based on the national production figures including:

- production of bread, cakes and biscuits,
- processed meat, fish, and poultry,
- sugar production (until 2004),
- production of margarine and solid cooking fats,
- production of animal feed,
- production of coffee,
- production of wine (distinguish between red and white),
- total production of beer,



- total production of spirits (other than Whisky and Brandy).

### Emission factors/ Emissions

NMVOC emissions from the food and beverage industry have been calculated using Tier 2 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.5.2.1.

**Table 4.5.2.1 Emission factors used for calculation of NMVOC emissions from food and drink**

	Value	Unit	Source: GB 2016, NFR 2.H.2
<b>Bread</b>	4.5	kg/t	Table 11 - Bread (typical) Europe
<b>Cakes and biscuits</b>	1	kg/t	Table 18
<b>Meat, fish, and poultry</b>	0.3	kg/t	Table 19
<b>Sugar</b>	10	kg/t	Table 20
<b>Margarine</b>	10	kg/t	Table 21
<b>Animal feed</b>	1	kg/t	Table 22
<b>Coffee roasting</b>	0.55	kg/t	Table 23
<b>Wine - red</b>	0,08	kg/hl	Table 25
<b>Wine - white</b>	0,035	kg/hl	Table 26
<b>Beer</b>	0,035	kg/hl	Table 27
<b>Spirits</b>	0.4	kg/hl alcohol	Table 32 – other spirits

### Recalculations

NMVOC emissions for the whole period have been recalculated because emissions from spirits have been excluded by mistake in the previous submission.

### Future improvements

No improvements are planned for this category.

## 4.6 Other production and consumption (NFR 2.I – 2.L)

Emission sources covered in this chapters are:

- 2I Wood processing
- 2K Consumption of POPs and heavy metals (e.g. electrical and scientific equipment)

Emissions from 2J Production of POPs and 2L Other production, consumption, storage, transportation or handling of bulk products do not occur in Slovenia and notation key NO has been used.

#### 4.6.1 Wood processing

NFR Code 2I

The present chapter addresses emissions of dust from the processing of wood. This includes manufacture of plywood, reconstituted wood products and engineered wood products. This source category is important for particulate emissions only.

##### Emission factors

Emissions of PM<sub>2.5</sub>, PM<sub>10</sub> and TSP from wood production have been taken from REMIS database.

##### Recalculations

No recalculations have been performed in this category.

##### Future improvements

No improvements are planned for this category.

#### 4.6.2 Consumption of POPs and heavy metals (e.g. electrical and scientific equipment)

NFR Code 2K

Production of electrical equipment containing PCB (transformers and capacitors) in Slovenia was terminated in January 1985. A study "A Concept of Handling the PCB/PCT in Slovenia" was made in 1999. PCB containing equipment has to be registered to Slovenian environment Agency - competent authority. It is also obligatory for the proprietors / owners of the PCB equipment to report to the competent authority, whether, when and how the PCB equipment was disposed of and where it was sent according to the principles of shipment of hazardous waste.

Electrical equipment, containing PCB in Slovenia:

- capacitor
- transformer

##### Methodology

To estimate emissions from consumption of POPs, the following methodology has been adopted:

$$E_{\text{pollutant}} = AR_{\text{production}} \times EF_{\text{pollutant}}$$

$E_{\text{pollutant}}$  – the emission of the specified pollutant

$AR_{\text{production}}$  – the activity rate for the production

$EF_{\text{pollutant}}$  – the emission factor for this pollutant

##### Activity data

Activity data for PCB emission calculations are obtained from Slovenian Environment Agency, Waste sector.

### Emission factors

PCB emissions from the electrical equipment have been calculated using Tier 3 emission factor from the relevant chapter of EMEP/EEA air pollutant emission inventory guidebook 2016, as presented in Table 4.6.2.1.

**Table 4.6.2.1 Emission factors used for calculation of PCB emissions from Consumption of POPs and heavy metals – electrical equipment**

	Value	Unit	Source: GB 2016, NFR 2.K
Capacitor	1.6	kg/t	Table 3-4
Transformer	0.06	kg/t	Table 3-4

### Recalculations

No recalculations have been performed in this category.

### Future improvements

No improvements are planned for this category.

## 5 AGRICULTURE

This chapter considers the emissions from manure management, application of inorganic N-fertilizers, animal manure and sewage sludge applied to soils, urine and dung deposited by grazing animals and farm-level agricultural operations including storage, handling and transport of agricultural products.

### 5.1 Manure management (3. B)

Sectors covered in this chapter are:

NFR Codes:

3B1a	Manure management - Dairy cattle
3B1b	Manure management - Non-dairy cattle
3B2	Manure management - Sheep
3B3	Manure management - Swine
3B4d	Manure management - Goats
3B4e	Manure management - Horses
3B4gi	Manure management - Laying hens
3B4gii	Manure management - Broilers
3B4giii	Manure management - Turkeys
3B4giv	Manure management - Other poultry
3B4h	Manure management - Other animals

#### Introduction

Ammonia (NH<sub>3</sub>) emissions which arise from excreta of farm animals are by far the most important source of ammonia emissions in Slovenia. It contributes 40 % of total emissions. High emissions are not only due to high emission factors, which are characteristic for animal production, but also due to specific structure of Slovenian agriculture. As a consequence of fact that about two thirds of utilized agricultural area is covered by grasslands, relatively high animal population, especially cattle, is maintained. Excreta of farm animals contribute also to emissions of nitric oxides (NO<sub>x</sub>), non-methane volatile organic compounds (NMVOC) and particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP). They contributed 0,2 %, 19,4 %, 0,9 %, 2,6% and 6,6 % of total NO<sub>x</sub>, NMVOCs, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP emissions, respectively.

This chapter considers the emissions of ammonia, nitric oxide and NMVOCs and particulate matter from animal housing and manure storage. Description of calculation procedure for application of manures and grazing animals is also a part of this chapter. However, emissions due to grazing and application of animal manures are reported under Crop production and agricultural soils chapter (NRF sector 3D).

#### Ammonia and nitric oxide

##### Methodology

The detailed (Tier 2) approach suggested by EMEP/EEA emission inventory guidebook, 2016 was used to assess the emissions of ammonia and nitric oxide. The methodology is based on principles of total ammonia nitrogen (TAN) fluxes through the manure management system. The model starts out with TAN excretions followed by emissions of NH<sub>3</sub>, N<sub>2</sub>O, NO and N<sub>2</sub> from animal

housing and manure stores. It was taken into account that only the nitrogen that was not lost from animal houses and manure stores is retained in animal manures. Therefore, emissions at each stage depend on the extent of emissions during the preceding stages. In case of slurry based systems mineralization of non -TAN N was taken into account and in the case of farmyard manure it was taken into account that a part of TAN is immobilised into organic matter.

### Activity data

The majority of activity data were obtained from the Statistical Office of the Republic of Slovenia (SORS). Data from 1991 are available on the SI-STAT data portal, under Environment and natural resources:

<http://pxweb.stat.si/pxweb/Database/Environment/Environment.asp>.

Data include the number of cattle, pigs, sheep, goats, horses, poultry and rabbits as well as average milk production per cow. Data for 1990 were obtained from old printed version of statistical yearbook. Data for some sub-categories of domestic animal species are missing for the certain years before the year 2000. Animals were distributed to these sub-categories based on the proportions in nearest years for which the data are available. For the rabbits no information on their number is available before the year 1997. Rounded value for 1997 was used for this period. There is also no information on the numbers of turkeys, ducks and geese for the period before 2000. These animals were treated in the frame of broilers for this period.

**Table 5.1.1 Number of farm animals in thousands**

Animal category	1990	1991	1992	1993	1994	1995	1996	1997	1998
<b>Cattle - total</b>	<b>532,9</b>	<b>483,9</b>	<b>503,8</b>	<b>477,5</b>	<b>477,4</b>	<b>495,5</b>	<b>486,2</b>	<b>445,7</b>	<b>453,1</b>
Dairy cows	225,3	205,7	213,0	203,7	197,4	197,1	154,7	147,6	146,5
Suckling cows	0,0	5,0	6,0	8,0	10,0	15,2	32,0	35,0	34,7
Other cattle	307,6	273,2	284,8	265,9	270,0	283,2	299,5	263,1	271,9
<b>Pigs - total</b>	<b>587,8</b>	<b>529,0</b>	<b>601,8</b>	<b>591,5</b>	<b>570,8</b>	<b>592,0</b>	<b>552,3</b>	<b>578,2</b>	<b>592,4</b>
Sows	57,7	51,9	55,5	55,1	55,9	56,2	47,9	52,8	52,2
Other breeding pigs*	10,7	9,3	10,6	10,4	9,9	9,9	10,2	11,6	10,1
Piglets	134,1	136,5	165,9	161,2	161,6	178,4	159,0	170,3	174,8
Fattening pigs**	385,4	331,4	369,9	364,8	343,5	347,5	335,2	343,4	355,2
<b>Small ruminants</b>	<b>30,2</b>	<b>38,5</b>	<b>32,0</b>	<b>37,2</b>	<b>39,8</b>	<b>51,1</b>	<b>55,8</b>	<b>65,8</b>	<b>89,2</b>
Sheep - total	20,3	28,5	22,0	26,6	29,1	39,1	43,2	51,9	72,4
Ewes	11,6	12,7	13,5	15,9	19,6	23,1	28,1	32,8	46,0
Other sheep	2,7	9,1	1,4	1,8	1,6	2,7	2,6	3,2	4,2
Lambs	6,0	6,7	7,1	8,9	7,9	13,3	12,5	15,9	22,2
Goats	10,0	10,0	9,9	10,6	10,7	11,9	12,6	13,9	16,8
Breeding female goats	6,7	6,7	6,7	6,9	7,8	8,3	9,5	10,2	11,4
Other goats	1,3	1,3	1,3	1,5	1,2	1,5	1,3	1,5	1,9
Kids	2,0	2,0	2,0	2,2	1,8	2,2	1,9	2,2	3,5
<b>Horses</b>	<b>10,4</b>	<b>10,8</b>	<b>8,9</b>	<b>8,5</b>	<b>8,1</b>	<b>8,0</b>	<b>8,5</b>	<b>9,9</b>	<b>12,1</b>
<b>Poultry - total</b>	<b>9753,2</b>	<b>10034,4</b>	<b>8734,0</b>	<b>6192,0</b>	<b>5794,0</b>	<b>4920,0</b>	<b>5573,0</b>	<b>7057,6</b>	<b>6407,1</b>
Laying hens	2340,5	2440,3	2323,0	1858,0	1840,0	1653,0	1615,0	1773,0	1695,2
Broilers	7412,7	7594,0	6411,0	4334,0	3954,0	3267,0	3958,0	5284,6	4711,9
Other chickens	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0
Turkeys	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0
Geese	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0
Ducks	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0
Other poultry	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0	0,0
<b>Rabbits-total</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>181,0</b>	<b>180,8</b>
Does	31,0	31,0	31,0	31,0	31,0	31,0	31,0	31,0	29,9
Other rabbits	150,0	150,0	150,0	150,0	150,0	150,0	150,0	150,0	150,8

\* Boars, gilts not yet covered

\*\* Including young breeding pigs

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(continued)

Animal category	1999	2000	2001	2002	2003	2004	2005	2006	2007
<b>Cattle - total</b>	<b>471,4</b>	<b>493,7</b>	<b>477,1</b>	<b>473,2</b>	<b>450,2</b>	<b>451,1</b>	<b>452,5</b>	<b>454,0</b>	<b>479,6</b>
Dairy cows	149,1	140,2	135,8	140,0	130,7	134,0	120,3	112,5	116,4
Suckling cows	36,5	53,9	52,8	55,0	55,2	48,1	57,0	60,5	61,2
Other cattle	285,8	299,5	288,5	278,3	264,4	269,1	275,3	281,0	301,9
<b>Pigs - total</b>	<b>558,5</b>	<b>603,6</b>	<b>599,9</b>	<b>655,7</b>	<b>620,5</b>	<b>534,0</b>	<b>547,4</b>	<b>575,1</b>	<b>542,6</b>
Sows	51,2	57,0	55,6	57,6	55,8	47,3	47,3	48,0	42,1
Other breeding pigs*	8,8	10,5	10,5	8,2	8,5	6,9	7,2	5,7	6,4
Piglets	161,8	178,3	181,2	179,0	182,2	158,0	159,4	161,6	154,0
Fattening pigs**	336,6	357,8	352,5	410,9	374,0	321,7	333,6	359,9	340,1
<b>Small ruminants</b>	<b>87,2</b>	<b>118,3</b>	<b>114,0</b>	<b>129,4</b>	<b>129,0</b>	<b>142,3</b>	<b>154,8</b>	<b>159,3</b>	<b>159,4</b>
Sheep - total	72,5	96,2	94,1	107,4	105,7	119,3	129,4	131,5	131,2
Ewes	50,8	66,3	66,0	75,9	72,1	84,4	89,7	89,1	90,8
Other sheep	3,4	5,3	5,1	5,3	4,9	5,3	5,5	6,2	6,2
Lambs	18,3	24,6	22,9	26,2	28,7	29,6	34,1	36,2	34,2
Goats	14,6	22,0	19,9	22,0	23,3	23,0	25,5	27,8	28,2
Breeding female goats	11,4	16,1	14,8	16,7	17,0	16,1	17,8	20,2	19,0
Other goats	1,3	2,4	2,3	2,1	2,1	2,1	2,4	2,7	2,6
Kids	1,9	3,6	2,8	3,1	4,2	4,9	5,3	4,9	6,6
<b>Horses</b>	<b>14,3</b>	<b>14,4</b>	<b>15,2</b>	<b>16,1</b>	<b>16,9</b>	<b>16,9</b>	<b>19,2</b>	<b>19,2</b>	<b>19,6</b>
<b>Poultry - total</b>	<b>5756,5</b>	<b>5105,9</b>	<b>5216,7</b>	<b>5265,7</b>	<b>4533,7</b>	<b>3268,0</b>	<b>3176,9</b>	<b>3056,7</b>	<b>4558,8</b>
Laying hens	1617,3	1539,5	1404,6	1401,1	1248,6	999,6	1085,3	1119,7	1338,4
Broilers	4139,2	2759,9	2879,9	2919,8	2523,8	1753,6	1598,5	1566,7	2837,4
Other chickens	0,0	483,0	589,4	446,4	503,7	336,5	312,1	232,4	177,9
Turkeys	0,0	252,1	251,0	417,3	209,3	130,2	135,4	110,1	158,0
Geese	0,0	2,5	4,0	3,3	3,1	3,5	3,4	1,9	2,6
Ducks	0,0	14,9	17,6	23,7	20,2	20,0	14,3	12,5	13,7
Other poultry	0,0	54,0	70,1	54,2	25,0	24,7	28,0	13,3	30,7
<b>Rabbits-total</b>	<b>180,5</b>	<b>180,3</b>	<b>166,5</b>	<b>152,7</b>	<b>139,0</b>	<b>134,5</b>	<b>130,1</b>	<b>122,8</b>	<b>115,6</b>
Does	28,8	27,7	27,0	26,3	25,6	24,7	23,8	23,0	22,2
Other rabbits	151,7	152,5	139,5	126,4	113,3	109,8	106,3	99,8	93,4

\* Boars, gilts not yet covered

\*\* Including young breeding pigs

(continued)

Animal category	2008	2009	2010	2011	2012	2013	2014	2015	2016
<b>Cattle - total</b>	<b>470,0</b>	<b>472,9</b>	<b>470,2</b>	<b>462,3</b>	<b>460,1</b>	<b>460,6</b>	<b>468,3</b>	<b>484,2</b>	<b>488,6</b>
Dairy cows	113,4	113,1	109,5	109,1	111,0	109,6	107,8	112,8	107,8
Suckling cows	62,6	61,0	63,9	61,7	56,5	56,2	60,5	57,0	63,5
Other cattle	294,0	298,8	296,8	291,6	292,5	294,8	299,9	314,3	317,3
<b>Pigs - total</b>	<b>432,0</b>	<b>415,2</b>	<b>395,6</b>	<b>347,3</b>	<b>296,1</b>	<b>288,4</b>	<b>281,3</b>	<b>271,4</b>	<b>265,7</b>
Sows	36,3	33,6	29,6	25,5	20,3	20,1	18,6	18,1	17,2
Other breeding pigs*	6,8	5,8	5,4	4,3	4,1	3,6	3,1	3,0	3,0
Piglets	121,7	108,6	99,0	81,6	66,0	67,5	63,6	59,5	57,5
Fattening pigs**	267,2	267,2	261,6	235,9	205,7	197,1	196,1	190,7	188,1
<b>Small ruminants</b>	<b>163,2</b>	<b>168,0</b>	<b>156,0</b>	<b>146,6</b>	<b>140,5</b>	<b>130,0</b>	<b>135,1</b>	<b>136,4</b>	<b>142,3</b>
Sheep - total	139,0	138,1	129,8	120,0	114,2	108,8	113,6	109,4	119,8
Ewes	95,0	95,2	90,9	81,5	77,3	73,4	78,0	75,2	81,5
Other sheep	6,7	7,3	6,4	6,1	6,0	5,5	5,4	5,5	6,9
Lambs	37,3	35,5	32,5	32,4	30,9	29,8	30,2	28,7	31,5
Goats	24,2	29,9	26,2	26,6	26,4	21,2	21,4	27,0	22,4
Breeding female goats	16,8	21,9	19,4	19,1	16,8	14,7	15,2	18,4	14,7
Other goats	2,4	2,8	2,4	2,6	2,8	2,1	2,2	2,5	2,3
Kids	5,0	5,3	4,4	4,9	6,8	4,5	4,0	6,1	5,4
<b>Horses</b>	<b>19,6</b>	<b>19,6</b>	<b>22,7</b>	<b>22,7</b>	<b>22,7</b>	<b>21,8</b>	<b>21,8</b>	<b>21,8</b>	<b>19,5</b>
<b>Poultry - total</b>	<b>4575,3</b>	<b>5211,9</b>	<b>4618,2</b>	<b>4006,7</b>	<b>4839,4</b>	<b>4907,0</b>	<b>5258,6</b>	<b>5753,9</b>	<b>6115,8</b>
Laying hens	1377,8	1553,2	1504,0	1365,2	1145,5	1380,0	1358,1	1458,1	1717,5
Broilers	2392,7	2944,6	2528,8	2154,8	3171,9	2827,2	3280,9	3479,2	3639,3
Other chickens	616,9	590,5	480,1	349,0	377,0	576,0	476,1	668,7	567,7

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Turkeys	144,6	94,5	68,9	95,8	110,9	96,2	121,4	108,1	156,2
Geese	2,9	2,7	2,1	1,9	2,2	2,8	1,7	3,1	3,4
Ducks	11,6	9,9	10,1	12,2	10,5	10,5	9,9	14,2	14,5
Other poultry	29,0	16,4	24,2	27,8	21,4	14,2	10,5	22,6	17,3
<b>Rabbits-total</b>	<b>105,4</b>	<b>95,2</b>	<b>85,1</b>	<b>88,8</b>	<b>92,5</b>	<b>96,2</b>	<b>100,7</b>	<b>105,2</b>	<b>109,8</b>
Does	20,6	19,0	17,4	18,6	19,8	21,0	22,3	23,6	24,9
Other rabbits	84,8	76,3	67,7	70,2	72,7	75,2	78,4	81,6	84,8

\* Boars, gilts not yet covered

\*\* Including young breeding pigs

(continued)

Animal category	2017
<b>Cattle - total</b>	<b>479,6</b>
Dairy cows	108,8
Suckling cows	59,9
Other cattle	310,9
<b>Pigs - total</b>	<b>257,2</b>
Sows	20,6
Other breeding pigs*	3,1
Piglets	58,7
Fattening pigs**	177,3
<b>Small ruminants</b>	<b>134,0</b>
Sheep - total	108,8
Ewes	77,9
Other sheep	5,0
Lambs	25,9
Goats	25,2
Breeding female goats	18,1
Other goats	2,6
Kids	4,5
<b>Horses</b>	<b>19,5</b>
<b>Poultry - total</b>	<b>6410,1</b>
Laying hens	1764,2
Broilers	3866,0
Other chickens	594,0
Turkeys	147,6
Geese	2,9
Ducks	17,4
Other poultry	18,0
<b>Rabbits-total</b>	<b>109,8</b>
Does	24,9
Other rabbits	84,8

\* Boars, gilts not yet covered

\*\* Including young breeding pigs

## Emission factors

In the first step nitrogen excretion from farm animals was estimated. It was obtained by multiplying the number of farm animals and nitrogen excretion rates on the level of individual animal species and categories. The nitrogen excretion rates, which were taken into account, are presented in Table 5.1.2. In dairy cows the nitrogen excretion has been linked to productivity, i.e. milk production (M). The equation proposed by Menzi et al. (1997) was used:

$$\text{N excretion (kg/year)} = 52,5 + 0,0105 \times \text{M (kg/year)} \quad (\text{eq. 1})$$

**Table 5.1.2 Nitrogen excretion rates for the calculation of ammonia emissions from animal production**

Animal category	N excretion (kg/year)	Source
<b>Cattle</b>		
Dairy cows	81-116	Equation 1
Suckling cows	78	Equation 1, taken into account 2400 kg of milk per year
Calves, fattening cattle, heifers	35	Menzi et al. (1997)
<b>Pigs</b>		
Sows <sup>a</sup>	36	EMEP/CORINAIR (2002)
Fattening pigs	14	EMEP/CORINAIR (2002)
<b>Small ruminants</b>		
Sheep <sup>b</sup>	15,5	EMEP/EEA (2016)
Goats <sup>c</sup>	15,5	EMEP/EEA (2016)
<b>Horses</b>	47,5	EMEP/EEA (2013)
<b>Poultry</b>		
Laying hens	0,71	Menzi et al. (1997)
Broilers	0,40	Menzi et al. (1997)
Turkeys	1,50	Döhler et al. (2002)
Geese	0,73	Döhler et al. (2002)
Ducks	0,60	Döhler et al. (2002)
<b>Rabbits<sup>d</sup></b>	8,1	IPCC (2006)

<sup>a</sup> Sows and pregnant gilts; the value includes N excretion in piglets and boars

<sup>b</sup> Adult sheep (including breeding female sheep and other adult sheep, like rams and barren sheep); the excretion value includes N excretion in lambs

<sup>c</sup> Adult goats (including breeding female goats and other adult goats, like he goats and barren goats); the excretion value includes N excretion in kids

<sup>d</sup> The excretion value applies for does; the value includes excretion in other rabbit categories

In case of dairy cows, where the N excretion was related to productivity, the value ranged from 81,6 to 115,8 kg of N per cow and year. Milk production and nitrogen excretion rates are presented in Table 5.1.3.

**Table 5.1.3 Milk production and nitrogen excretion (Nex) rates for dairy cattle in kg/head/year**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
<b>Milk production (kg/year)</b>	2775	3252	2835	2800	3014	3170	3831	3975	4091	4252
<b>Nex (kg N per animal per year)</b>	81,6	86,6	82,3	81,9	84,1	85,8	92,7	94,2	95,5	97,1
	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
<b>Milk production (kg/year)</b>	4625	4807	5198	5062	4853	5479	5708	5726	5764	5531
<b>Nex (kg N per animal per year)</b>	101,1	103,0	107,1	105,7	103,5	110,0	112,4	112,6	113,0	110,6
	2010	2011	2012	2013	2014	2015	2016	2017		
<b>Milk production (kg/year)</b>	5517	5516	5593	5435	5717	5598	6024	5954		
<b>Nex (kg N per animal per year)</b>	110,4	110,4	111,2	109,6	112,5	111,3	115,8	115,0		

In certain species of domestic animals, nitrogen excretions of some animal categories (mostly young animals like piglets, lambs and kids or male breeding animals like boars) are considered



to be covered by excretion factors of other categories, like sows, does, adult sheep or adult goats. As a result, average excretion rates reported in CRF differ from those given in Table 5.1.2. Average excretion rates which were calculated by dividing the total N excretion by total number of animals are given in Table 5.1.4. Due to variation in proportions of individual categories within animal species the average excretion rates differ slightly among years.

**Table 5.1.4 Average nitrogen excretion (N<sub>ex</sub>) rates for animal species in which nitrogen excretions of some animal categories are considered to be covered by other categories. The values refer to total population (kg N/head/year)**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
Pigs	12,7	12,3	11,9	12,0	11,9	11,6	11,6	11,6	11,6	11,7
Sheep	10,9	11,9	10,5	10,3	11,3	10,2	11,0	10,8	10,7	11,6
Goats	12,4	12,4	12,4	12,3	12,9	12,7	13,2	13,0	12,3	13,5
Rabbits	1,39	1,39	1,39	1,39	1,39	1,39	1,39	1,39	1,34	1,29
	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
Pigs	11,7	11,6	11,9	11,7	11,6	11,6	11,8	11,6	11,7	11,9
Sheep	11,5	11,7	11,7	11,3	11,7	11,4	11,2	11,5	11,3	11,5
Goats	13,0	13,3	13,3	12,7	12,2	12,3	12,8	11,9	12,3	12,8
Rabbits	1,25	1,32	1,40	1,49	1,49	1,48	1,52	1,56	1,58	1,61
	2010	2011	2012	2013	2014	2015	2016	2017		
Pigs	12,0	12,2	12,2	12,1	12,1	12,2	12,2	12,2		
Sheep	11,6	11,3	11,3	11,3	11,4	11,4	11,4	11,8		
Goats	12,9	12,6	11,5	12,2	12,6	12,0	11,7	12,8		
Rabbits	1,65	1,70	1,73	1,77	1,80	1,82	1,84	1,84		

### **Emissions from animal housing, manure stores and due to fertilization with animal manures in cattle production**

Emission factors, which tell us how much of N from animal excreta is lost to the atmosphere in the form of ammonia, depend on manure management systems. Factors, along with some basic information on manure management systems in cattle production, are presented in Table 5.1.7. Generally, EMEP/EEA factors were used. In case of introduction of abatement techniques the basic emission factors were multiplied by (1- efficiency coefficient). Efficiency coefficients were obtained either from EMEP/EEA manual or from Draft revised United Nations Economic Commission for Europe Framework Code for Good Agricultural Practice for Reducing Ammonia Emissions (ECE/EB AIR/2014/8). The fraction of individual manure management systems was estimated on the basis of the results of farm census data from 1991 and 2000. Since manure management systems were not reported in the census, data on size and structure of cattle-breeding farms were used for rough estimates. It was considered that all farms with less than 10 head of bovine animals had solid manure storage systems, that 30 % of farms with 10-19 head of animals practiced liquid manure storage and 70 % of them solid manure storage, and that all farms with 20 cows or more had liquid manure storage systems. Linear regression was used to estimate the changes in manure management systems in the period 1990-2000. After 2000, data on farm size and structure were reported by the Statistical Office for the years 2003, 2005, 2007, 2010, 2013 and 2016. For the years with missing values the proportions of various manure storage systems were obtained by interpolation or extrapolation. Animals kept in liquid systems were further divided into animals kept in liquid manure storage with natural crust cover, animals kept in liquid manure storage below animal confinements and animals from which the excreta was treated in anaerobic digesters. Based on information on manure management that was collected in the frame of milk recording service on a large number of dairy farms in 2005

(Babnik and Verbič, 2007) it was estimated that the ratio between slurry stored in stores with natural crust and slurry stored below animal confinements is 0,46:0,54. Based on information from the same source the solid manure was divided into farmyard manure stored in heaps and deep bedding (0,90:0,10). The proportion of slurry treated in anaerobic digesters was estimated on the basis of data collected from biogas plants by the means of interview (data provided by Poje, unpublished). Based on above mentioned data and data on total number of cattle it was estimated that during the period 2006-2010 the proportion of digested cattle manures increased from 0,03 to 0,36 %. Anaerobic digesters were not markedly spread thereafter and therefore the same value was used for the period 2011-2017.

The fraction of grazing bovine animals for 1990 has been estimated on the basis of data on grazing animals on mountain pastures and expert estimate on the scale of grazing on intensive grasslands (Verbič et al., 1999). In 2000, all grazing animals on mountain and other pastures were recorded. This census showed that in 2000, one way or another, 21 % of animals were grazing. This data have been corrected with regard to the length of the grazing season, considering the fact that animals on mountain pastures will graze for 141 days on the average, and on other pastures for 210 days. As result, the corrected proportion of grazed animals for 2000 was estimated to be 0,117. The same procedure was used for the data obtained by sample survey on agricultural production methods in 2010. It showed that the corrected proportion of grazed animals increased to 0,126.

The estimate for 1990 was used for the period 1985-1990. For the period 1991-1999, the data on grazing were obtained by linear regression which was calculated on the basis of data for the years 1990 and 2000 and for the period 2001-2009 the estimates obtained by linear regression for the years 2000 and 2010. For the years thereafter, extrapolated values based on 2000-2010 period were used. It has been estimated that the fraction of grazing animals and the fraction of liquid manure management systems have increased while the fraction of bovine animals in straw based systems has decreased. Detailed information on grazing and distribution of manure management systems is given in Table 5.1.6.

It has to be pointed out, that in case of farmyard manure system, one part of excreta is stored as solid (faeces + bedding) while the other part (urine + manure effluents) is stored as liquid. It was taken into account that cattle excrete 57 % of N in urine and 43 % in faeces. It is incorporated into calculation process. As a result, the proportion of manure storage systems in CRF is not equal to proportions of manure management systems reported in Table 5.1.6. An example is given in a Table 5.1.5.

**Table 5.1.5 Example of conversion of proportions of various animal rearing systems into proportions of manure storage systems**

Rearing system	Proportion	N distribution into storage systems	Storage system		
			Liquid	Solid	Grazing
Slurry	0,568	100 % liquid	0,568	0,000	0,000
Farmyard manure	0,303	57 % liquid 43 % solid	0,173	0,130	0,000
Grazing	0,129	100 % grazing	0,000	0,000	0,129
<b>Total</b>	<b>1,000</b>		<b>0,741</b>	<b>0,130</b>	<b>0,129</b>

**Table 5.1.6 Distribution of various manure management systems in cattle production. In farmyard manure system part of N is retained in solid and part in liquid fraction**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
<b>Grazing</b>										
Dairy cows	0,059	0,065	0,071	0,076	0,082	0,088	0,094	0,100	0,105	0,111
Other cows	0,059	0,065	0,071	0,076	0,082	0,088	0,094	0,100	0,105	0,111
Other cattle	0,066	0,071	0,076	0,081	0,086	0,092	0,097	0,102	0,107	0,112
<b>Farmyard manure</b>										
Dairy cows	0,593	0,579	0,565	0,551	0,537	0,523	0,509	0,495	0,481	0,467
Other cows	0,593	0,579	0,565	0,551	0,537	0,523	0,509	0,495	0,481	0,467
Other cattle	0,588	0,575	0,561	0,548	0,534	0,521	0,507	0,494	0,480	0,467
<b>Slurry</b>										
Dairy cows	0,348	0,356	0,365	0,373	0,381	0,389	0,397	0,405	0,414	0,422
Other cows	0,348	0,356	0,365	0,373	0,381	0,389	0,397	0,405	0,414	0,422
Other cattle	0,346	0,354	0,362	0,371	0,379	0,388	0,396	0,405	0,413	0,422
	<b>2000</b>	<b>2001</b>	<b>2002</b>	<b>2003</b>	<b>2004</b>	<b>2005</b>	<b>2006</b>	<b>2007</b>	<b>2008</b>	<b>2009</b>
<b>Grazing</b>										
Dairy cows	0,117	0,118	0,119	0,120	0,121	0,122	0,122	0,123	0,124	0,125
Other cows	0,117	0,118	0,119	0,120	0,121	0,122	0,122	0,123	0,124	0,125
Other cattle	0,117	0,118	0,119	0,120	0,121	0,122	0,122	0,123	0,124	0,125
<b>Farmyard manure</b>										
Dairy cows	0,453	0,435	0,418	0,400	0,395	0,390	0,373	0,356	0,341	0,327
Other cows	0,453	0,435	0,418	0,400	0,395	0,390	0,373	0,356	0,341	0,327
Other cattle	0,453	0,435	0,418	0,400	0,395	0,390	0,373	0,356	0,341	0,327
<b>Slurry</b>										
Dairy cows	0,430	0,447	0,463	0,480	0,484	0,488	0,504	0,521	0,534	0,548
Other cows	0,430	0,447	0,463	0,480	0,484	0,488	0,504	0,521	0,534	0,548
Other cattle	0,430	0,447	0,463	0,480	0,484	0,488	0,504	0,521	0,534	0,548
	<b>2010</b>	<b>2011</b>	<b>2012</b>	<b>2013</b>	<b>2014</b>	<b>2015</b>	<b>2016</b>	<b>2017</b>		
<b>Grazing</b>										
Dairy cows	0,126	0,127	0,127	0,128	0,129	0,130	0,131	0,131		
Other cows	0,126	0,127	0,127	0,128	0,129	0,130	0,131	0,131		
Other cattle	0,126	0,127	0,127	0,128	0,129	0,130	0,131	0,131		
<b>Farmyard manure</b>										
Dairy cows	0,312	0,309	0,306	0,303	0,292	0,281	0,270	0,270		
Other cows	0,312	0,309	0,306	0,303	0,292	0,281	0,270	0,270		
Other cattle	0,312	0,309	0,306	0,303	0,292	0,281	0,270	0,270		
<b>Slurry</b>										
Dairy cows	0,562	0,564	0,567	0,569	0,579	0,589	0,599	0,599		
Other cows	0,562	0,564	0,567	0,569	0,579	0,589	0,599	0,599		
Other cattle	0,562	0,564	0,567	0,569	0,579	0,589	0,599	0,599		

**Table 5.1.7 Emission factors and basic information on manure management systems for the calculation of NH<sub>3</sub>, N<sub>2</sub>O, NO and N<sub>2</sub> emissions in cattle production (Sources for emission factors: Menzi et al., 1997, EMEP/EEA emission inventory guidebook, 2013, ECE/EB AIR/2014/8)**

	Grazing	Tied housing system		Loose housing system
		Farmyard* manure	Liquid* fraction (urine)	Slurry
<b>Proportion of TAN at the level of excretion (in kg/kg total N)*</b>	0,60	0,30	0,70	0,60
<b>Basic information</b>				
Proportion of covered manure stores	/	0,00	0,90	0,50
Proportion of manure application in favourable weather conditions or incorporation	/	0,20	0,20	0,20
Bedding material (kg per animal per year)	0	Cows: 730 kg Other cattle: 240 kg	0	0
N added in bedding (kg per animal per year)	0,00	Cows: 2,92 kg Other cattle: 0,96 kg	0,00	0,00
Mineralization of non-TAN N during storage (proportion of total non-TAN N)	/	0,00	0,00	0,10
Immobilization of TAN during storage (proportion of TAN)	/	0,0067	0,0000	0,0000
<b>Emission factors (kg NH<sub>3</sub>-N/kg TAN)</b>				
From animal houses or during grazing (proportion of excreted TAN)	Dairy cattle: 0,1 Other cattle: 0,06	0,190	0,200	0,200
Emissions from uncovered manure stores (proportion of TAN entering the stores)	/	0,270	0,200	0,200
Emissions from covered manure stores (proportion of TAN entering the stores)	/	/	0,040	0,040
Emissions due to manure application – basic coefficients (proportion of TAN leaving the stores)	/	0,790	0,550	0,550
Emissions due to manure application – coefficients for manure incorporation or application in favourable weather conditions (proportion of TAN leaving the stores)	/	0,474	0,330	0,330
<b>Emission factors (kg N<sub>2</sub>O-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	/	0,080	0,001	0,001
<b>Emission factors (kg NO-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	/	0,0080	0,0001	0,0001
<b>Emission factors (kg N<sub>2</sub>-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	/	0,300	0,003	0,003

\* in farmyard manure system it was taken into account that 0,57 of N was retained in solid and 0,43 in liquid fraction.

Based on expert estimate it was assumed for the entire reporting period that 20 % of animal manures was applied to fields in favourable weather conditions or incorporated into the soil within about 12 hours after application (Table 5.1.7). It was assumed that basic emission coefficients for the above mentioned practices are reduced by 40 % (ECE/EB AIR/2014/8). For the period 2015-2017 it was also taken into account that a certain part of slurry was applied by the means of low emission techniques (9,5, 11 and 11 % for arable land and 2,2, 2,2 and 2,2 %

for grasslands in years 2015, 2016 and 2017 respectively). The information is based on the area supported by Rural development programme (operation "low emission fertilization"). For the arable land it was considered that low emission techniques were equally distributed into trailing hoses, trailing shoe, shallow injection and deep injection. For grasslands trailing hoses were assumed. For the efficiency of low emission techniques the values proposed by ECE/EB AIR/2014/8 were taken into account.

### **Emissions from animal housing, manure stores and due to fertilization with animal manures in pig production**

To obtain reliable estimates on the manure management systems in pig production the population was disaggregated into three categories:

- a) commercial pig farms,
- b) market oriented family farms, and
- c) small scale family farms.

Data published by the SORS allow a breakdown of the entire herd into commercial pig farms and family farms for the period 1986-2002. Family farms were further divided into market oriented and small scale farms. In 1986, the estimate of production for market oriented family farms was based on the data on acquisition of pigs from market oriented family farm production, which was published by the SORS. The number of swine in small scale family farm production has been estimated from the difference between the entire herd and market oriented production (commercial and market oriented family farms). For 2000, the number of pigs in the small scale family farm production has been estimated on the basis of the census of agricultural holdings. Pigs kept on farms with up to 10 pigs have been considered as small scale family farm production, pigs on family farms which kept more than 10 pigs have been considered as market oriented family farm production. From 1986 to 2000, the fraction of pigs in small scale family farm production kept diminishing. In the period between 1986 and 2000, the proportion of small scale production was obtained by interpolation. After 2000, data on farm structure for the years 2003, 2005, 2007, 2010, 2013 and 2016 have been reported by the SORS. These data were used to estimate the number of pigs on small scale family farms. For the years with non-existing data on farm structure (2001, 2002, 2004, 2006, 2008, 2009, 2011, 2012, 2014, 2015) the numbers of pigs on small scale family farms were obtained by interpolating the values for neighbouring years. For 2017 the same proportion as for 2016 was used. For the period after the year 2002 the number of pigs on commercial farms could not be obtained directly from the data reported by SORS. Therefore, it was estimated using the data on farm structure for the years 2003, 2005, 2007, 2010, 2013 and 2016. The estimate is based on the number of pigs which are kept on farms with more than 399 pigs. The pigs belonging to this category (pigs kept on farms with more than 400 pigs) were allocated among commercial and market oriented family farms on the basis of their proportion in the year 2000. The pigs kept on farms with 10 to 399 pigs were entirely allocated to market oriented family farms. For the year 2017 the data on farm structure were not gathered. Therefore, the same structure as for 2016 was taken into account.

For market oriented family farm production, it was considered that 95 % of animal excreta were collected in the form of liquid manure and 5 % in the form of solid manure. For small scale family farm production, it was estimated that 95 % of pigs is reared in solid manure storage systems and 5 % in liquid manure systems. For the big commercial pig farms old-style separators were characteristic for the period 1985 to 1994. App. 20 % of solids was separated from liquid manure by the use of these separators. The remainder (80 %) was either treated in lagoons (75 %) or spread as liquid manure (25 %). The time from 1995 to 1999 was a period of introducing new separators and the beginning of operation of anaerobic digesters. Introducing new separators on commercial farms increased the estimated portion of separated solid phase to 40 %.

Detailed information on manure management systems are given in Table 5.1.8. Emission factors for pig production are given in Table 5.1.9.

**Table 5.1.8 Distribution of various manure management systems in pig production**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
Slurry	0,281	0,250	0,345	0,360	0,355	0,351	0,341	0,366	0,374	0,401
Farmyard manure	0,355	0,375	0,323	0,315	0,311	0,287	0,291	0,266	0,246	0,245
Separation (solid fraction)	0,091	0,094	0,083	0,081	0,084	0,197	0,200	0,201	0,207	0,238
Anaerobic lagoons	0,274	0,281	0,249	0,244	0,251	0,148	0,150	0,151	0,155	0,064
Anaerobic digestion	0,000	0,000	0,000	0,000	0,000	0,016	0,017	0,017	0,017	0,051
	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
Slurry	0,503	0,494	0,536	0,525	0,507	0,488	0,486	0,490	0,489	0,499
Farmyard manure	0,221	0,213	0,209	0,201	0,199	0,197	0,184	0,171	0,182	0,192
Separation (solid fraction)	0,187	0,198	0,173	0,185	0,199	0,212	0,159	0,153	0,127	0,128
Anaerobic lagoons	0,050	0,053	0,046	0,050	0,053	0,057	0,043	0,041	0,034	0,034
Anaerobic digestion	0,040	0,042	0,037	0,040	0,043	0,046	0,129	0,144	0,169	0,147
	2010	2011	2012	2013	2014	2015	2016	2017		
Slurry	0,541	0,547	0,554	0,560	0,553	0,545	0,538	0,538		
Farmyard manure	0,202	0,211	0,220	0,229	0,228	0,226	0,224	0,224		
Separation (solid fraction)	0,126	0,118	0,109	0,101	0,106	0,111	0,116	0,116		
Anaerobic lagoons	0,000	0,000	0,000	0,000	0,000	0,000	0,000	0,000		
Anaerobic digestion	0,131	0,124	0,117	0,109	0,114	0,118	0,122	0,122		

### **Emissions from animal housing, manure stores and due to fertilization with animal manures in poultry production**

Emissions in poultry production were calculated as a sum of emissions for broilers, layers, ducks, turkeys and geese. For broilers, turkeys, geese and ducks exclusively floor system on bedding was assumed. For laying hens, combined floor system (1/4) and battery-cage systems (3/4) were assumed for 1990. Assumption was made on the basis of expert estimate. It was also assumed that in 50 % the manure is removed daily and stored in tanks (liquid system) while in 50 % it is collected under the batteries (i.e. poultry manure without bedding). After introduction of dung drying system to certain farms, new estimates were obtained for 2002. Layers which were assumed to be kept in floor system, in system where manure is collected under the batteries and in dung drying system were allocated to solid system. Layers which were assumed to be kept in system where the manure is removed daily and stored in tanks was allocated to liquid systems. Emission factors for poultry rearing are given in Table 5.1.10.

**Table 5.1.9 Emission factors and basic information on manure management systems for the calculation of NH<sub>3</sub>, N<sub>2</sub>O, NO and N<sub>2</sub> emissions in pig production (Sources for emission factors: EMEP/EEA emission inventory guidebook, 2013, EPA, 2004)**

	Farmyard manure and solid*	Slurry	Anaerobic lagoon	Anaerobic fermenter
<b>Proportion of TAN at the level of excretion (in kg/kg total N)*</b>	0,70	0,70	0,70	0,70
<b>Basic information</b>				
Proportion of covered manure stores	0,00	0,50	0,00	1,00
Proportion of manure application in favourable weather conditions or immediate incorporation	0,20	0,20	/	0,20
Bedding material (kg per animal per year)	FP: 200 S: 600	0	0	0
N added in bedding (kg per animal per year)	FP: 0,8 S: 2,4	0	0	0
Mineralization of non-TAN N during storage (proportion of total non-TAN N)	0	0,1	1	0,1
Immobilization of TAN during storage (proportion of TAN)	0,0067	0,000	0,000	0,000
<b>Emission factors (kg NH<sub>3</sub>-N/kg N)</b>				
From animal houses (proportion of excreted TAN)	FP: 0,27 S: 0,25	FP: 0,28 S: 0,22	FP: 0,28 S: 0,22	FP: 0,28 S: 0,22
Emissions from uncovered manure stores (proportion of TAN entering the stores)	0,45	0,14	0,71	0,14
Emissions from covered manure stores (proportion of TAN entering the stores)	/	0,028	/	0,028
Emissions due to manure application – basic coefficients (proportion of TAN leaving the stores)	0,810	0,400	/	0,400
Emissions due to manure application – coefficients for immediate manure incorporation or application in favourable weather conditions (proportion of TAN leaving the stores)	0,486	0,240	/	0,240
<b>Emission factors (kg N<sub>2</sub>O-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	FYM: 0,05 Solid: 0,08	0,00	0,00	0,00
<b>Emission factors (kg NO-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	0,0080	0,0001	0,0001	0,0001
<b>Emission factors (kg N<sub>2</sub>-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	0,300	0,003	0,290	0,003

\* solid fraction extracted from slurry during the separation process

Abbreviations: FP – Fattening pigs, S – Sows, FYM – farmyard manure

For the low emission application techniques and their effectiveness the same assumptions as for cattle manures were used.

**Table 5.1.10 Emission factors for the calculation of NH<sub>3</sub>, N<sub>2</sub>O, NO and N<sub>2</sub> emissions in poultry production (Source for emission factors: EMEP/EEA emission inventory guidebook, 2013)**

	Laying hens - solid	Laying hens - liquid	Broilers	Ducks	Turkeys	Geese
<b>Proportion of TAN at the level of excretion (in kg/kg total N)*</b>	0,70	0,70	0,70	0,70	0,70	0,70
<b>Basic information</b>						
Proportion of manure application in favourable weather conditions or immediate incorporation	0,20	0,20	0,20	0,20	0,20	0,20
Bedding material (kg per animal per year)	0*	/	0*	0*	0*	0*
N added in bedding (kg per animal per year)	0*	/	0*	0*	0*	0*
Mineralization of non-TAN N during storage (proportion of total non-TAN N)	0,00	0,10	0,00	0,00	0,00	0,00
<b>Emission factors (kg NH<sub>3</sub>-N/kg N)</b>						
From animal houses (proportion of excreted TAN)	0,41	0,41	0,28	0,24	0,35	0,57
Emissions from manure stores (proportion of TAN entering the stores)	0,14	0,14	0,17	0,24	0,24	0,16
Emissions due to manure application – basic coefficients (proportion of TAN leaving the stores)	0,690	0,690	0,660	0,540	0,540	0,450
Emissions due to manure application – coefficients for immediate manure incorporation or application in favourable weather conditions (proportion of TAN leaving the stores)	0,414	0,414	0,396	0,324	0,324	0,270
<b>Emission factors (kg N<sub>2</sub>O-N/kg TAN)</b>						
Emissions from manure stores (proportion of TAN entering the stores)	0,040	0,000	0,030	0,030	0,030	0,030
<b>Emission factors (kg NO-N/kg TAN)</b>						
Emissions from manure stores (proportion of TAN entering the stores)	0,008	0,0001	0,008	0,008	0,008	0,008
<b>Emission factors (kg N<sub>2</sub>-N/kg TAN)</b>						
Emissions from manure stores (proportion of TAN entering the stores)	0,30	0,003	0,30	0,30	0,30	0,30

\* Sawdust; considered to contain no available N and to have no TAN immobilization potential

### **Emissions from animal housing, manure stores and due to fertilization with animal manures in small ruminants, horses and rabbits**

Ammonia emissions in goats, sheep, horses and rabbits were estimated using the information presented in Table 5.1.11. The proportions of grazing animals were estimated by the means of expert opinion. It was estimated that during the grazing season all sheep, 80 % of goats and 50



% of horses were grazed. Two hundred and fifty days of grazing season has been considered for sheep and 210 for goats and horses. For the remaining period it has been considered that these animals were kept in straw based systems. It was considered that rabbits are not grazed.

**Table 5.1.11 Emission factors and basic information on manure management systems for the calculation of NH<sub>3</sub>, N<sub>2</sub>O, NO and N<sub>2</sub> emissions in sheep, goats, horses and rabbits (Source for emission factors: EMEP/EEA emission inventory guidebook, 2013)**

	Sheep	Goats	Horses	Rabbits
<b>Proportion of TAN at the level of excretion (in kg/kg total N)*</b>	0,50	0,50	0,60	0,50 <sup>a</sup>
<b>Basic information</b>				
Proportion of manure application in favourable weather conditions or immediate incorporation	0,20	0,20	0,20	0,20
Bedding material (kg per animal per year)	91	91	1460	3,65
N added in bedding (kg per animal per year)	0,365	0,365	5,84	0,015
Immobilization of TAN during storage (proportion of TAN)	0,0067	0,0067	0,0067	0,0067
<b>Emission factors (kg NH<sub>3</sub>-N/kg N)</b>				
From animal houses (proportion of excreted TAN)	0,22	0,22	0,22	0,22 <sup>a</sup>
During grazing (proportion of excreted TAN)	0,09	0,09	0,35	/
Emissions from manure stores (proportion of TAN entering the stores)	0,280	0,280	0,350	0,280 <sup>a</sup>
Emissions due to manure application – basic coefficients (proportion of TAN leaving the stores)	0,090	0,090	0,090	0,090
Emissions due to manure application – coefficients for immediate manure incorporation or application in favourable weather conditions (proportion of TAN leaving the stores)	0,054	0,054	0,054	0,054
<b>Emission factors (kg N<sub>2</sub>O-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	0,070	0,070	0,080	0,080 <sup>b</sup>
<b>Emission factors (kg NO-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	0,008	0,008	0,008	0,008
<b>Emission factors (kg N<sub>2</sub>-N/kg TAN)</b>				
Emissions from manure stores (proportion of TAN entering the stores)	0,30	0,30	0,30	0,30

<sup>a</sup> There are no emission factors in EMEP/EEA emission inventory guidebook; values for sheep were used

<sup>b</sup> There are no emission factors in EMEP/EEA emission inventory guidebook; value for horses were used

## **Non-methane volatile organic compounds (NMVOCs)**

### **Methodology**

With exception of rabbits, the detailed (Tier 2) approach suggested by EMEP/EEA emission inventory guidebook, 2013 was used to assess the emissions of NMVOCs. For cattle the methodology based on gross energy intake and for other animal species methodology based on excretion of volatile substance was used. Total NMVOC emissions were estimated as a sum of emissions from silage stores, from the silage feeding, from housing, from manure stores, from manure application and from grazing. Country specific data for gross energy intake were used

to estimate emissions in cattle production. The information was obtained from national UNFCCC reporting. Based on information that high dry matter grass and maize silages which are characterised by low concentrations of volatile fatty acids are produced in Slovenia (Verbič et al., 2011) the suggested emission factors for silage storage and feeding (EMEP/EEA emission inventory guidebook, 2013) were reduced correspondingly. For rabbits, default emission factor which was suggested by EMEP/EEA (2016) was used.

### Activity data

The activity data were obtained from the SORS. They include the number of cattle, pigs, sheep, goats, horses, poultry and rabbits.

### Emission factors

#### *Emissions in cattle production*

Emissions in cattle were estimated on the basis of gross energy intake which was reported to UNFCCC. The gross energy intake depends on several factors among which the most important are milk production in dairy cows and growth rate in fattening cattle. As a result of increased productivity the estimated gross energy intake in dairy cows and other cattle increased considerably during the period 1985 – 2013 (Table 5.1.12). The fraction of silage in diet was estimated on the basis of survey which was performed in 2005 (Verbič et al., 2006) and the fact that silage making in Slovenia became an important forage preservation method after the year 1970. For the period 1985 - 2004 the proportions of silage in diet was obtained by interpolation of data taken into account that there was no silage in the diets in the year 1970 and that its proportion in 2005 was 0,55. The estimate for 2005 was used also for the period after 2005. For the proportion of time spent on grazing the same data was used as for emissions of ammonia and nitric oxide.

Emission factors for calculation of NMVOC emissions are given in Table 5.1.12. The emissions from silage stores were calculated by multiplying the values for silage feeding by a fixed value of 0,25 as suggested by EMEP/EEA emission inventory guidebook, 2013. The emissions from manure stores and emissions due to manure application were also estimated indirectly on the basis of emissions from animal houses. It was supposed that the relation between NMVOC emissions from animal houses on the one hand and emissions from manure stores and application of manure on the other is the same as for ammonia.

**Table 5.1.12 Emission factors and basic information used for calculation of NMVOC emissions in cattle (Source for emission factors: EMEP/EEA emission inventory guidebook, 2013)**

	Dairy cows	Suckling cows	Other cattle
<b>Basic information</b>			
Gross energy intake (MJ yr <sup>-1</sup> per animal)	78549 - 106309	73752-74272	40408 - 44309
Time spent in animal houses (proportion of total)	0,869 – 0,941	0,869 – 0,941	0,869 – 0,934
Fraction of silage in diet (proportion of maximal possible dry matter quantity in the diet)	0,31 – 0,55	0,31 – 0,55	0,31 – 0,55
The share of the emission in silage store compared to the emission from the feeding table	0,25	0,25	0,25
<b>Emission factors</b>			
Emissions due to silage feeding (kg NMVOC MJ <sup>-1</sup> gross energy intake from silage)*	0,0001201	0,0001201	0,0001201

Emissions from housing (kg NMVOC MJ <sup>-1</sup> gross energy intake in animal houses)	0,0000353	0,0000353	0,0000353
Emissions from grazing (kg NMVOC MJ <sup>-1</sup> gross energy intake during grazing)	0,0000069	0,0000069	0,0000069

\* EF which was suggested by EMEP/EEA emission inventory guidebook, 2013 was reduced by 40 % due to high dry matter silages which are characterised by restricted fermentation.

### *Emissions in pigs, sheep, goats, horses, poultry and rabbits*

Emissions in small ruminants, horses, pigs and poultry were estimated on the basis of volatile solids excretion using the same values as reported to UNFCCC (i.e. default values according to IPCC, 2006). It was assumed that no silage is given to these animals. For the proportion of time spent on grazing the same data was used as for emissions of ammonia and nitric oxide.

The emissions from animal houses and from grazing areas were calculated on the basis of emission factors which are given in Table 5.1.13. The emissions from manure stores and emissions due to manure application were also estimated indirectly on the basis of emissions from animal houses. It was supposed that the relation between NMVOC emissions from animal houses on the one hand and emissions from manure stores and application of manure on the other is the same as for ammonia. For rabbits, a default EMEP/EEA (2016) emission factor was used (0,059 kg per animal and year).

**Table 5.1.13 Emission factors and basic information used for calculation of NMVOC emissions in cattle (Source for emission factors: EMEP/EEA emission inventory guidebook, 2013)**

	Volatile solids (VS) (kg yr <sup>-1</sup> per animal)	Time spent in animal houses (proportion of total)	EF housing (kg NMVOC kg <sup>-1</sup> VS excreted)	EF grazing (kg NMVOC kg <sup>-1</sup> VS excreted)
Sheep	146	0,315	0,0016140	0,00002349
Goats	110	0,540	0,0016140	0,00002349
Horses	777	0,712	0,0016140	0,00002349
Fattening pigs	110	1,000	0,0017030	/
Sows	168	1,000	0,0070420	/
Layers	7,30	1,000	0,0056840	/
Broilers	3,65	1,000	0,0091470	/
Turkeys	25,55	1,000	0,0056840	/

## **Particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>, TSP)**

### **Methodology**

The methodology suggested by EMEP/EEA emission inventory guidebook, 2016 was used to assess the emissions of particulate matter. Due to opinion that a scientific literature as a whole does not support the use of Tier 2 methodology (EMEP/EEA, 2016) it was decided to use a Tier 1 approach.

### **Activity data**

The activity data were obtained from the SORS. They include the number of cattle, pigs, sheep, goats, horses and poultry. For cattle, pigs and poultry the emissions were estimated on the level of subcategories.

## Emission factors

Emission factors are presented in Table 5.1.14. They apply to housed animals only. The number of housed animals was calculated by multiplying the total number of animals by the fraction of housed animals. The latest was obtained from information on proportion of grazing animals as described in methodology which was used for calculation of emissions of ammonia and nitric oxide.

**Table 5.1.14 Emission factors used for calculation of TSP, PM<sub>10</sub> and PM<sub>2.5</sub> emissions from livestock husbandry (housing) (Source: EMEP/EEA emission inventory guidebook, 2016)**

Livestock	TSP (kg/head)	PM <sub>10</sub> (kg/head)	PM <sub>2.5</sub> (kg/head)
Dairy cattle	1,38	0,63	0,41
Non-dairy cattle (including young cattle, beef cattle and suckling cows)	0,59	0,27	0,18
Non-dairy cattle (calves)	0,34	0,16	0,1
Sheep <sup>a</sup>	0,14	0,06	0,02
Pigs (fattening pigs)	1,05	0,14	0,006
Pigs (weaners)	0,27	0,05	0,002
Pigs (sows)	0,62	0,17	0,01
Goats <sup>b</sup>	0,14	0,06	0,02
Horses	0,48	0,22	0,14
Laying hens <sup>c</sup>	0,19	0,04	0,003
Broilers	0,04	0,02	0,002
Other poultry (chickens)	0,04	0,02	0,002
Turkeys	0,11	0,11	0,02
Ducks	0,14	0,14	0,02
Geese	0,24	0,24	0,03
Other poultry	0,04	0,02	0,002

<sup>a</sup> adult sheep, including barren sheep and rams

<sup>b</sup> adult goats, including barren goats and he goats

<sup>c</sup> including parents of broilers

There is no information whether emission factors of particulate matter include or exclude condensable component.

## Recalculations

No recalculations were done since last reporting.

## Future improvements

No further improvements are planned until the next submission.

## Manure management - Buffalo: NFR Code 3B4a

## Manure management - Mules and asses: NFR Code 3B4f

Notation Key "NO" (not occurring) was used for these sectors, since no additional livestock exist within a country. No emissions originate from these sectors.

## 5.2 Crop production and agricultural soils (3. D)

Sectors covered in this chapter are:

NFR Codes:

3Da1	Inorganic N-fertilizers (includes also urea application)
3Da2a	Animal manure applied to soils
3Da2b	Sewage sludge applied to soils
3Da3	Urine and dung deposited by grazing animals
3Dc	Farm-level agricultural operations including storage, handling and transport of agricultural products

Agricultural soils are source of ammonia (NH<sub>3</sub>), nitric oxides (NO<sub>x</sub>), non-methane volatile organic compounds (NMVOCs) and particulate matter. They contributed 50,6 %, 6,6 % and 0,1 % of total NH<sub>3</sub>, NO<sub>x</sub> and NMVOCs emissions, respectively. Contribution of PM<sub>2.5</sub> and PM<sub>10</sub> were 0,1 % and 1,3 %. The main sources of ammonia are application of inorganic N-fertilizers and nitrogen which is excreted by grazed farm animals. Small quantities of ammonia are emitted also due to application of sewage sludge. Four sources of NO emissions from agricultural soils were identified, i.e. application of synthetic N-fertilizers, application of animal manures, nitrogen deposited to soils by grazed farm animals and application of sewage sludge, the latest being almost negligible. Crop production is also source of particulate matter while NMVOCs are emitted due to animal grazing.

### 5.2.1 Inorganic N-fertilizers

NFR Code 3Da1

#### Ammonia

##### Methodology

Ammonia emissions due to use mineral fertilizers were assessed according to EMEP/EEA emission inventory guidebook, 2016 methodology. They were obtained by multiplying data on consumption of nitrogen from mineral fertilizers and emission factors for three main groups of fertilizers.

##### Activity data

The consumption of nitrogen from mineral fertilizers in agriculture has been obtained from Statistical office of the republic of Slovenia (SORS). There is a sharp increase in sales of mineral fertilizers observed in 1992. The reasons for increase of activity data and consequently strong increase in NH<sub>3</sub> emission between 1991 and 1992 are:

- poor economic situation and war for independence in 1991 which causes considerable lower sales of mineral fertilizers than during the previous years,
- independence and improved economic situation in 1992,
- high inflation in 1992 which stimulated farmers to renew stocks of mineral fertilizers (well established practice from the times of high inflation in Yugoslavia was to invest in material resources),

- main supplier of mineral fertilizers in Slovenia was (and it still is) a company from Croatia. The fear that due to political situation in Croatia there will be a disturbance in mineral fertilizers supply forced farmers to increase stocks of mineral fertilizers.

**Table 5.2.1.1 Consumption of mineral fertilizers according to fertilizer type (in tonnes of N)**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
Total	27169	23758	38938	33376	33944	32235	31296	33999	34801	34380
CAN	10866	9477	15491	13242	13467	12269	12576	13338	13716	13545
Urea	5437	4805	7957	6891	7010	7697	6145	7323	7369	7290
NP, NPK	10866	9477	15491	13242	13467	12269	12576	13338	13716	13545
	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
Total	34159	34765	33412	34501	30264	29169	30383	29613	25039	28202
CAN	13365	13607	12639	13204	11757	10930	11715	11506	10197	9873
Urea	7429	7552	8134	8094	6749	7309	6954	6600	4645	8456
NP, NPK	13365	13607	12639	13204	11757	10930	11715	11506	10197	9873
	2010	2011	2012	2013	2014	2015	2016	2017		
Total	27486	27134	26300	27263	28612	28319	27097	27084		
CAN	10261	10551	9624	10386	11350	11417	10582	10938		
Urea	6964	6032	7051	6492	5911	5485	5932	5207		
NP, NPK	10261	10551	9624	10386	11350	11417	10582	10938		

### Emission factors

Emission factors 0,008, 0,155 and 0,050 kg NH<sub>3</sub>-N per kg of N were used for calcium ammonium nitrate (CAN), urea and other mineral (NP and NPK) fertilizers respectively. Data for urea consumption for the period 1994-2016 were obtained from SORS (personal communication, data not officially published). For the period 1985-1993 the proportion of urea in total mineral-N fertilizer consumption was estimated by extrapolation based on 1994-2013 period. The allocation of the rest of mineral-N fertilizes between CAN and other (NP and NPK) fertilizers were done on the basis of expert judgement (50:50). Fertilizers which are characterized by high emission factors are not in use (anhydrous ammonia) or even prohibited (ammonium carbonate fertilizers). For the years 2016 and 2017 it was taken into account that low emission application techniques are used on 8,8 % and 11,8 % of arable land, respectively. It was considered that 60 % of urea is used on arable land and that urea incorporation reduces ammonia emissions by 50 %. The decision was made on the basis of the fact that investments in machinery which enables urea incorporation are supported by the Rural development programme.

### Recalculations

Error in calculation of ammonia emissions from urea application was discovered in previous report (factor 0,015 was used instead of 0,155). The value was corrected. As a result, ammonia emissions for the entire reporting period increased.

### Future improvements

No further improvements are planned until the next submission.

## **Nitric oxide**

### **Methodology**

Nitric oxide emissions due to use mineral fertilizers were assessed according to EMEP/EEA emission inventory guidebook, 2016 methodology. No Tier 2 methodology is available and therefore Tier 1 methodology was used. The emissions were obtained by multiplying data on consumption of nitrogen from mineral fertilizers and emission factor.

### **Activity data**

The consumption of nitrogen from mineral fertilizers in agriculture has been obtained from the SORS.

### **Emission factors**

An uniform emission factor, i.e. 0,040 kg NO<sub>2</sub> per kg of N applied in form of synthetic fertilizers, was used (EMEP/EEA emission inventory guidebook, 2016).

### **Recalculations**

On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result the emissions of nitric oxide were reduced for the entire reporting period.

### **Future improvements**

No further improvements are planned until the next submission.

## **5.2.2 Animal manure applied to soils**

NFR sector 3Da2a

## **Ammonia**

Emissions of ammonia following the application of animal manure are reported under this chapter. Calculation methods are presented in the frame of chapter Manure management (3B).

## **Nitric oxide**

### **Methodology**

Nitric oxide which is released from soils due to fertilization with animal manures is reported under this chapter. Emissions were assessed according to EMEP/EEA emission inventory guidebook, 2016 methodology. No Tier 2 methodology is available and therefore Tier 1 methodology was used. Emissions were obtained on the basis of data on nitrogen which is returned to soil by the means of animal manures and adequate emission factor.

### **Activity data**

Data on nitrogen which is returned to soil in form of animal manures were calculated within methodology described in chapter Manure management (NFR sector 3B).

### **Emission factors**

An emission factor 0,040 kg NO<sub>2</sub> per kg of nitrogen which is applied to soil in form of animal manures was used (EMEP/EEA emission inventory guidebook, 2016).

### **Recalculations**

The ammonia emissions for 2015 and 2016 were corrected by taking into account the new information on the low emission manure spreading techniques which were supported by Rural development programme. As a result, the emissions of ammonia for the above mentioned years were reduced.

On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced for the entire reporting period.

### **Future improvements**

No further improvements are planned until the next submission.

## **5.2.3 Sewage sludge applied to soils**

NFR Code 3Da2b

### **Ammonia**

#### **Methodology**

Default emission factor, as suggested by EMEP/EEA emission inventory guidebook (2016) was used.

#### **Activity data**

Since 2000, data on sewage sludge application to the agricultural soils have been obtained from the reports prepared under the Sewage sludge directive (Environment Agency of the Republic of Slovenia). Data for 1995 and 1998 were obtained from environmental reports. It was assumed that the same proportion of sewage sludge (30 %) have been deposited to agricultural land for the period before 1995. Data for 1996, 1997 and 1999 were estimated by interpolation. Due to rigorous restrictions the application of sewage sludge to agricultural land is extremely small.



**Table 5.2.3.1 Application of sewage sludge to agricultural soils (in tonnes of N)**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
Sewage sludge	78	78	78	78	78	78	70	62	55	33
	2000	2001	2002	2003	2004	2005	2006	2007	2008	2009
Sewage sludge	12	20	43	18	5	3	1	1	0,4	0,4
	2010	2011	2012	2013	2014	2015	2016	2017		
Sewage sludge	18	0,04	0,04	0,04	7,18	0,51	18,31	0,00		

**Emission factors**

An emission factor 0,13 kg of ammonia nitrogen per kg of total ammonia nitrogen applied by sewage sludge was used (EMEP/EEA 2016). It was taken into account that 0,70 of total sewage sludge nitrogen is in the form of ammonia (data for solid pig manure; EMEP/EEA emission inventory guidebook, 2013). For the nitrogen content in sewage sludge the value 3,9 % (on dry matter basis) was used.

**Recalculations**

The emission factor 0,81 kg of ammonia nitrogen per kg of total ammonia nitrogen applied by sewage sludge, which was used in previous report, was replaced by a factor 0,13 kg of ammonia nitrogen per kg of total ammonia nitrogen applied by sewage sludge (EMEP/EEA 2016). It resulted in a decrease of ammonia emissions.

**Future improvements**

No further improvements are planned until the next submission.

**Nitric oxide**

Emissions of nitric oxide following the application of sewage sludge are more or less negligible. It can happen that the use of sewage sludge in agriculture will increase in future and therefore the source was not neglected.

**Methodology**

The Tier 1 approach suggested by EMEP/EEA 2016 emission inventory guidebook was used to assess the emissions of nitric oxide.

**Activity data**

Data sources on sewage sludge application to the agricultural soils are described in the frame of ammonia methodology (see text above).

**Emission factors**

An emission factor 0,040 kg NO<sub>2</sub> per kg of nitrogen which is applied to soil in form of sewage sludge was used as suggested by EMEP/EEA emission inventory guidebook (2016).

## **Recalculations**

Recalculations for the entire period were done. On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced.

## **Future improvements**

No further improvements are planned until the next submission.

### **5.2.4 Other organic fertilizers applied to soils**

NFR Code 3Da2c

Emissions due to application of other organic fertilizers to soils were not reported in previous submissions. It was considered that the quantities of compost applied to soils were negligible. TERT recommended that the use of compost should be analysed with the aim to get activity data until the next submission. Slovenia started the activities to get the data on quantities of compost and its composition from producers. However, due to reporting dynamics data are not ready yet.

### **5.2.5 Urine and dung deposited by grazing animals**

NFR sector 3Da3

## **Ammonia**

### **Introduction**

Ammonia emissions due to nitrogen in animal excreta deposited during grazing is minor source of ammonia emissions. They contribute less than 2 % of total emissions.

### **Methodology**

Ammonia emissions due to N excretion on pasture were calculated within methodology described in chapter Manure management (NFR sector 3B). The emissions are reported under this chapter.

### **Activity data**

For activity data regarding the emissions due to nitrogen in animal excreta deposited during grazing see chapter on Manure management (NFR sector 3B).

### **Emission factors**

Emission factors used for calculation of the emissions due to nitrogen in animal excreta deposited during grazing are given in chapter on Manure management (NFR sector 3B) (Tables 5.1.7, 5.1.9 and 5.1.11).

### **Recalculations**

No recalculations were done since the previous report.

### **Future improvements**

No further improvements are planned until the next submission.

### **Nitric oxide**

#### **Methodology**

Nitric oxide emissions due to nitrogen deposited to agricultural soils by grazing animals were assessed according to EMEP/EEA emission inventory guidebook, 2016 methodology. No Tier 2 methodology is available and therefore Tier 1 methodology was used. Emissions were obtained by multiplying the amount of nitrogen returned to soils by grazed farm animals by an adequate emission factor.

#### **Activity data**

Data on nitrogen which is returned to soil by grazed farm animals were calculated within methodology described in chapter Manure management (NFR sector 3B).

#### **Emission factors**

An emission factor 0,040 kg NO<sub>2</sub> per kg of N returned to soils by grazed farm animals was used (EMEP/EEA emission inventory guidebook, 2016).

#### **Recalculations**

Recalculations for the entire period were done. On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced.

#### **Future improvements**

No further improvements are planned until the next submission.

### **Non-methane volatile organic compounds (NMVOCs)**

#### **Methodology**

NMVOCs emissions due grazing were calculated within methodology described in chapter Manure management (NFR sector 3B). The emissions are reported under this chapter.

### **Activity data**

For activity data regarding the emissions due to grazing see chapter on Manure management (NFR sector 3B).

### **Emission factors**

Emission factors used for calculation of the emissions due to grazing are given in chapter on Manure management (NFR sector 3B) (Tables 5.1.7, 5.1.9 and 5.1.11).

### **Recalculations**

No recalculations were performed since last submission.

### **Future improvements**

No further improvements are planned until the next submission.

## **5.2.6 Farm-level agricultural operations including storage, handling and transport of agricultural products**

NFR Code 3Dc

### **Particulate matter (PM<sub>2.5</sub>, PM<sub>10</sub>)**

#### **Methodology**

The detailed (Tier 2) approach suggested by EMEP/EEA emission inventory guidebook, 2016 was used to assess the emissions of particulate matter from crop production. Emissions from soil cultivation, harvesting, cleaning and drying of crops were estimated.

#### **Activity data**

The activity data were obtained from the SORS. They include the areas of arable land as well as temporary and permanent grasslands. Some cereals which are characterised by a specific emission factors (wheat and spelt, rye and triticale, barley, oat) were treated separately.

#### **Emission factors**

Emission factors for PM<sub>10</sub> and PM<sub>2.5</sub> are presented in Tables 5.2.6.1 and 5.2.6.2. These factors refer to wet climate conditions. With the exemption of grasslands it was considered that each operation is carried out once a year. In case of temporary grasslands it was considered that cultivation appears once per two years only. It was also considered that 30 % of grasslands (temporary and permanent) is harvested as a hay and that harvesting is carried out twice a year. The areas of crop types which were used for assessment of PM<sub>10</sub> and PM<sub>2.5</sub> are presented in Table 5.2.6.3.

**Table 5.2.6.1 Emission factors used for calculation of PM<sub>10</sub> emissions from crop production (Source: EMEP/EEA emission inventory guidebook, 2016)**

Crop	Soil cultivation (kg/ha per year)	Harvesting (kg/ha per year)	Cleaning (kg/ha per year)	Drying (kg/ha per year)
Wheat (including spelt)	0,25	0,49	0,19	0,56
Rye (including triticale)	0,25	0,37	0,16	0,37
Barley	0,25	0,41	0,16	0,43
Oat	0,25	0,62	0,25	0,66
Other arable	0,25	NC	NC	NC
Temporary grasslands	0,125 <sup>a</sup>	0,15 <sup>b</sup>	0	0
Permanent grasslands	0	0,15 <sup>b</sup>	0	0

<sup>a</sup> given that permanent grasslands are cultivated once per two years (estimate) EMEP/EEA (2016) factor (0,25 kg/ha per operation) was divided by two

<sup>b</sup> factor based on estimate that 30% of meadows are harvested as a hay and that hay making is performed twice a year. EMEP/EEA (2016) factor (0,25 kg/ha per operation) was multiplied by 0,3 and 2 (0,25×0,3×2=0,15).

**Table 5.2.6.2 Emission factors used for calculation of PM<sub>2.5</sub> emissions from crop production (Source: EMEP/EEA emission inventory guidebook, 2016)**

Crop	Soil cultivation (kg/ha per year)	Harvesting (kg/ha per year)	Cleaning (kg/ha per year)	Drying (kg/ha per year)
Wheat (including spelt)	0,015	0,02	0,009	0,168
Rye (including triticale)	0,015	0,015	0,008	0,111
Barley	0,015	0,016	0,008	0,129
Oat	0,015	0,025	0,0125	0,198
Other arable	0,015	NC	NC	NC
Temporary grasslands	0,0075 <sup>a</sup>	0,006 <sup>b</sup>	0	0
Permanent grasslands	0	0,006 <sup>b</sup>	0	0

<sup>a</sup> given that permanent grasslands are cultivated once per two years (estimate) EMEP/EEA (2016) factor (0,015 kg/ha per operation) was divided by two

<sup>b</sup> factor based on estimate that 30% of meadows are harvested as a hay and that hay making is performed twice a year. EMEP/EEA (2016) factor (0,01 kg/ha per operation) was multiplied by 0,3 and 2 (0,01×0,3×2=0,006)

The PM<sub>2.5</sub> and PM<sub>10</sub> emission factors represent filterable PM emissions.

**Table 5.2.6.3 Areas of various crop types in Slovenia in 000 ha**

	1990	1991	1992	1993	1994	1995	1996	1997	1998	1999
Wheat (including spelt)	43,50	39,43	36,41	37,17	35,88	36,78	35,16	33,43	35,03	31,62
Rye (including triticale)	2,63	2,74	2,69	2,64	2,10	2,29	2,28	1,78	1,71	1,55
Barley	7,49	7,86	8,15	9,09	12,65	12,72	12,54	10,83	10,87	10,94
Oat	2,74	2,37	2,38	2,39	2,59	1,87	1,89	1,82	1,79	2,41
Other arable	162,35	178,42	172,38	173,40	171,16	166,36	163,89	155,90	155,87	154,55

2019 INFORMATIVE INVENTORY REPORT for SLOVENIA

Temporary grasslands	28,38	23,99	23,58	23,21	21,31	24,68	21,63	21,06	20,37	20,86
Permanent grasslands	310,37	334,33	333,30	330,36	319,11	308,67	300,81	289,99	287,47	296,59
	<b>2000</b>	<b>2001</b>	<b>2002</b>	<b>2003</b>	<b>2004</b>	<b>2005</b>	<b>2006</b>	<b>2007</b>	<b>2008</b>	<b>2009</b>
Wheat (including spelt)	38,26	39,34	35,73	35,59	32,39	30,06	32,08	32,04	35,41	34,53
Rye (including triticale)	1,51	1,97	2,28	2,45	3,23	3,31	3,64	3,91	3,96	4,29
Barley	11,57	12,66	12,39	13,79	15,32	15,45	17,04	18,53	19,23	20,09
Oat	2,25	1,92	2,01	1,96	1,85	2,73	2,47	2,33	1,89	1,77
Other arable	157,52	152,50	149,93	157,82	153,61	156,49	154,00	151,66	152,60	148,27
Temporary grasslands	16,76	23,63	24,03	24,19	27,65	27,70	29,21	30,22	33,93	36,48
Permanent grasslands	308,20	307,04	307,18	308,35	286,83	304,91	285,00	297,28	285,97	267,30
	<b>2010</b>	<b>2011</b>	<b>2012</b>	<b>2013</b>	<b>2014</b>	<b>2015</b>	<b>2016</b>	<b>2017</b>		
Wheat (including spelt)	31,95	29,67	34,59	31,76	33,12	30,73	31,46	28,02		
Rye (including triticale)	4,27	4,16	4,54	4,98	5,87	5,73	6,26	6,11		
Barley	18,73	17,48	17,97	17,31	18,48	20,11	19,18	20,37		
Oat	1,77	1,84	1,37	1,20	1,36	1,51	1,33	1,45		
Other arable	143,26	142,71	147,96	149,20	150,84	147,57	150,79	151,86		
Temporary grasslands	35,50	34,30	32,96	34,42	32,56	30,37	31,78	32,48		
Permanent grasslands	285,71	262,60	281,16	277,48	279,92	278,68	276,25	279,22		

### Recalculations

It was found that for 1990 the calculation procedures were not properly linked to data on the areas of arable land, temporary and permanent grasslands. As a result emissions from soil cultivation, harvesting, cleaning and drying of crops were not estimated. Error was eliminated. It did not affect the results for the period 1991-2016.

### Source-specific planned improvements

No improvements are planned for this source.

### 5.2.7 Field burning of agricultural residues

NFR Code: 3F

Burning of agricultural residues is banned. It has also not been practiced before the ban. The main reason is shortage of bedding material. About two thirds of total agricultural area is covered by grasslands. In addition, a lot of forage crops are produced on arable land. Cereals cover only about 13 % of total agricultural area and a demand on the local market is high. The price of straw (up to 0.2 € per kg in 2018) is close to price of cereal grains. Maize stover and other residues which are not used for bedding is incorporated into soil. Notation Key "NO" (not occurring) was used for this activity.

**Other organic fertilizers applied to soils: NFR Code 3Da2c**

**Crop residues applied to soils: NFR Code 3Da4**

**Indirect emissions from managed soils: NFR Code 3Db**

**Off-farm storage, handling and transport of bulk agricultural products: NFR Code 3Dd**

**Cultivated crops: NFR Code 3De**

**Use of pesticides: NFR Code 3Df**

**Field burning of agricultural residues: NFR Code 3F**

**Agriculture other: NFR Code 3I**

Notation Key "NO" (not occurring) was used for these sectors, since no activity or process exist within a country. No emissions originate from these sectors.

## 6 WASTE

This chapter covers emissions resulting from solid waste disposal on land, from treatment of liquid wastes and waste incineration. Waste management and treatment of industrial and municipal wastes are minor sources of air pollutant emissions.

Sectors covered in this chapter are:

NFR Codes:

5A	Biological treatment of waste - Solid waste disposal on land
5B1	Biological treatment of waste - Composting
5C1a	Municipal waste incineration
5C1bii	Hazardous waste incineration
5C1biii	Clinical waste incineration
5C1bv	Cremation
5D1	Domestic wastewater handling
5D2	Industrial wastewater handling
5E	Other waste

### 6.1 Biological treatment of waste - Solid waste disposal on land

NFR Code 5A

#### Introduction

This chapter treats emissions from solid waste disposal on land. This source is only a minor source of air pollutant emissions. Major emissions from waste disposal are emissions of greenhouse gases, predominantly CH<sub>4</sub>.

#### Methodology

To estimate emissions of NMVOC from waste disposal the following methodology has been adopted:

$$E = q \times EF$$

E – emission (g)

q – quantity of total waste disposed (t)

EF – emission factor (g/t)

#### Activity data

For calculation of NMVOC and particulate matter emissions from solid waste disposal on land, the relevant activity data is total amount of waste disposed at municipal solid waste disposal sites.

Detailed description on activity data used for calculation is presented in National Inventory Report 2017, chapter CH<sub>4</sub> Emissions from Solid Waste Disposal sites, pg 283.  
[http://unfccc.int/national\\_reports/annex\\_i\\_ghg\\_inventories/national\\_inventories\\_submissions/it](http://unfccc.int/national_reports/annex_i_ghg_inventories/national_inventories_submissions/it)



ems/10116.php (Slovenia, NIR, SVN NIR 2017.pdf)

Quantities of landfilled waste in the period 1990-2017 are presented in Table 6.1.1.

**Table 6.1.1 Quantity of total waste disposed**

Year	Waste disposed (t)	Year	Waste disposed (t)
1990	671980	2004	727464
1991	681580	2005	752546
1992	687897	2006	840338
1993	694418	2007	811674
1994	702108	2008	822722
1995	707000	2009	750743
1996	725000	2010	623224
1997	743000	2011	504997
1998	761000	2012	387421
1999	780000	2013	274724
2000	800000	2014	257914
2001	820000	2015	260828
2002	821436	2016	113280
2003	820132	2017	142622

### Emission factors

A default emission factors for NMVOC, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP were used for emissions calculation. Emission factors were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 5A Biological treatment of waste - Solid waste disposal on land, Table 3-1, pg. 5.

**Table 6.1.2 Emission factors for solid waste disposal on land**

Pollutant	Value	Unit	References
NMVOC	1,56	kg/t	Emission Inventory Guidebook, 2016, 5A Biological treatment of waste - Solid waste disposal on land, Table 3-1, pg. 5
PM <sub>2.5</sub>	0,033	g/t	Emission Inventory Guidebook, 2016, 5A Biological treatment of waste - Solid waste disposal on land, Table 3-1, pg. 5
PM <sub>10</sub>	0,219	g/t	Emission Inventory Guidebook, 2016, 5A Biological treatment of waste - Solid waste disposal on land, Table 3-1, pg. 5
TSP	0,463	g/t	Emission Inventory Guidebook, 2016, 5A Biological treatment of waste - Solid waste disposal on land, Table 3-1, pg. 5

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Very small quantities of NMVOC and particulates are emitted from solid waste disposal on land. The contribution of this activity to the total NMVOC is 0,6 %. Emissions of particulate matter are negligible.

NMVOC emissions are dependent on total annual amount of municipal waste and the fraction of landfilled municipal waste. The quantities of municipal waste have marked a decrease in

recent years. Possible explanations is that the quantities in previous years have mostly been arrived at by estimation, whereas in the last four years we had at our disposal very accurate data from all solid waste disposal sites. At the same time, the area where waste is collected separately and then recycled is getting ever wider. NMVOC, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP emissions for the period 1990-2017 are presented in Figures 6.1.1 - 6.1.4.

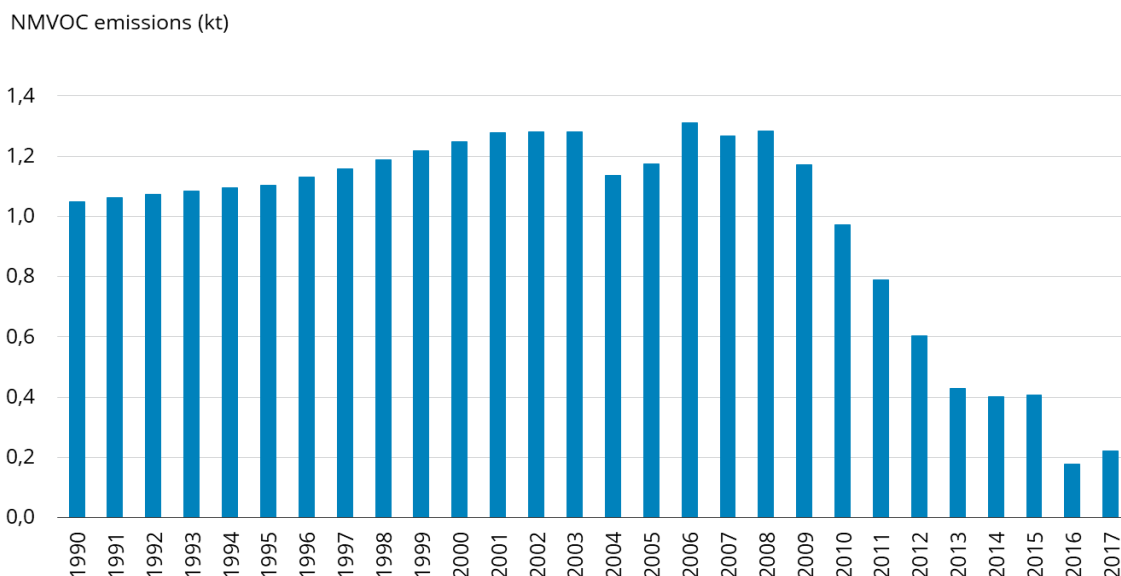


Figure 6.1.1 NMVOC emissions from solid waste disposal on land

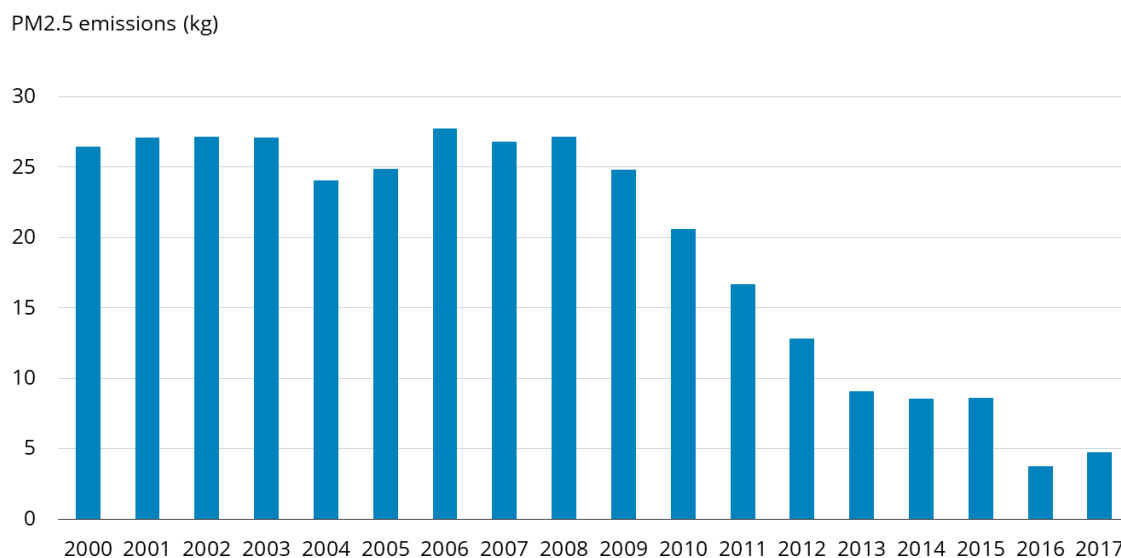
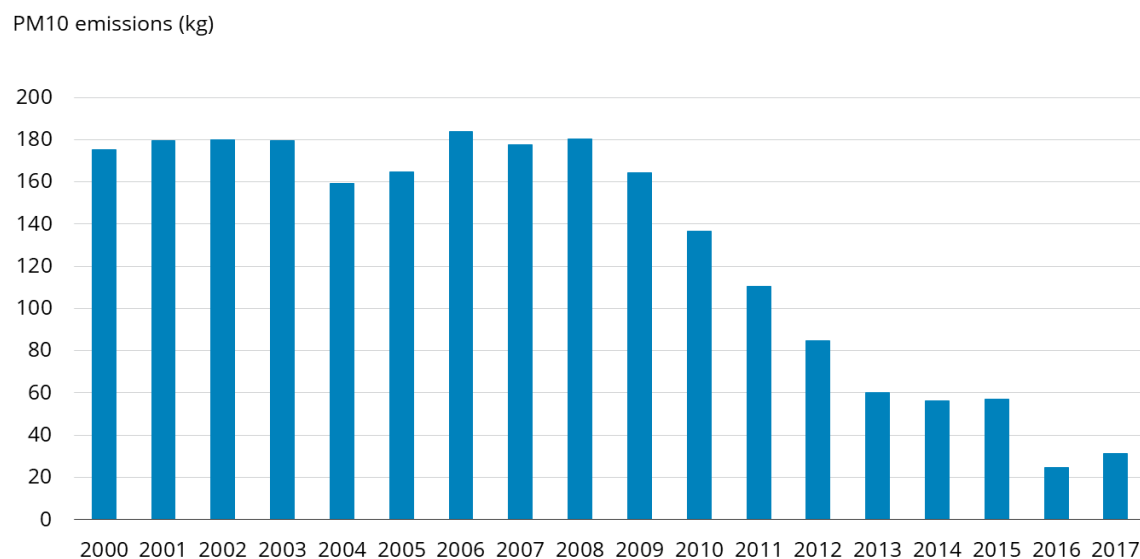
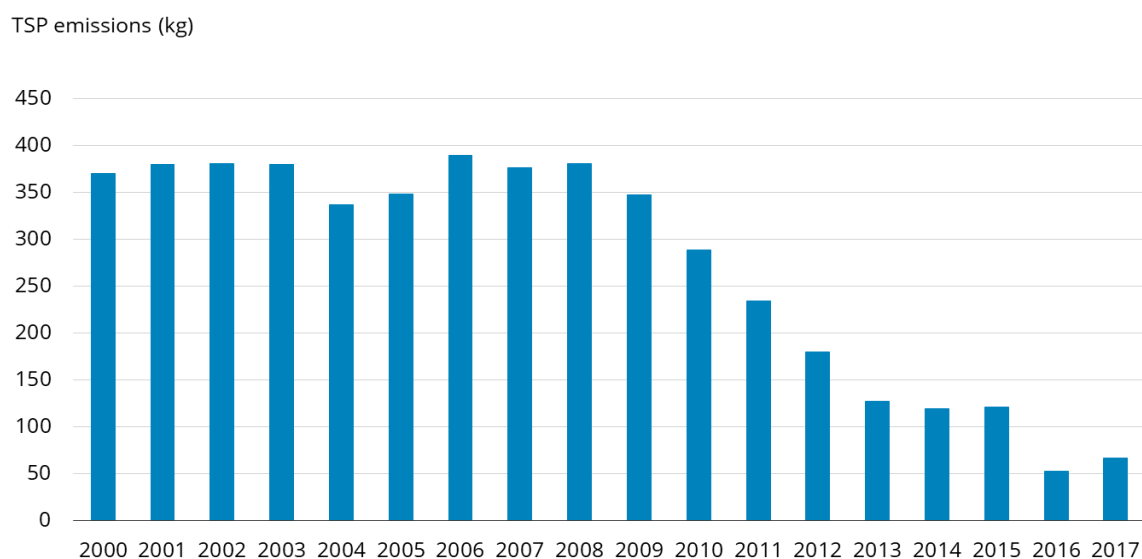


Figure 6.1.2 PM<sub>2.5</sub> emissions from solid waste disposal on land



**Figure 6.1.3 PM<sub>10</sub> emissions from solid waste disposal on land**



**Figure 6.1.4 TSP emissions from solid waste disposal on land**

### Recalculations

Recalculations of NMVOC, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP were recalculated for the year 2016 due to new data on amount of landfilled waste obtained.

### Category-specific QA/QC and verification

Amount of solid waste disposed on land have been thoroughly examined. Data obtained from

Statistical Office of the Republic of Slovenia was used for emission calculation. Emission factors applied were checked as well. According to 2017 in-depth EU NECD review 2017 recommendation EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 was used for emission calculations.

### Future Improvements

No improvement is planned for this category.

## 6.2 Biological treatment of waste – Composting

NFR Code 5B1

### Introduction

This chapter covers the emissions from the biological treatment of waste – composting. This source is not significant on a national level for any pollutant, only a small amount of ammonia is produced.

### Methodology

To estimate emissions of NH<sub>3</sub> from waste composting the following methodology has been adopted:

$$E = q \times EF$$

E – emission (g)

q – quantity of waste composted (t)

EF – emission factor (g/t)

### Activity data

For calculation of NH<sub>3</sub> emissions from composting the relevant activity data is an annual amount of total organic waste composted in wet weight. Activity data were obtained from Statistical Office of the Republic of Slovenia for the period 2002-2017. Data for the period 1995-2001 were estimated due to unavailability of precise annual data for years before 2002. There was no composting prior the year 1995.

**Table 6.2.1 Quantity of organic waste composted**

Year	Waste composted (t)	NH <sub>3</sub> emissions (t)
1995-2001	31542	7,57
2002	31542	7,57
2003	31803	7,63
2004	23367	5,61
2005	14930	3,58
2006	11537	2,77
2007	14867	3,57
2008	18196	4,37

<b>2009</b>	22896	5,50
<b>2010</b>	26671	6,40
<b>2011</b>	49763	11,94
<b>2012</b>	49000	11,76
<b>2013</b>	66215	15,89
<b>2014</b>	70395	16,89
<b>2015</b>	72366	17,37
<b>2016</b>	74355	17,85
<b>2017</b>	97860	23,48

### Emission factors

Emission factor for NH<sub>3</sub> was taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016, 5.B.1 Biological treatment of waste - composting, compost production, Table 3-1, pg. 5. The value for NH<sub>3</sub> emission factor is 0,24 kg/t organic waste.

### Emissions

Very small quantities of NH<sub>3</sub> are emitted from composting. The contribution of this activity to the total NH<sub>3</sub> emissions in the year 2017 is below 0,1 %. Emissions for the period 1995-2017 are presented in the Table 6.2.1.

### Recalculations

No recalculations were performed since last submission.

### Future Improvements

No improvement is planned for this category.

## 6.3 Municipal waste incineration

NFR Code 5C1a

### Introduction

This sector includes emissions from domestic and commercial refuse, often referred to as 'municipal solid waste' (MSW). Municipal solid waste is the unwanted material collected from households and commercial organisations. It consists of a mix of combustible and non-combustible materials, such as paper, plastics, food waste, organic waste from home gardens, glass, defunct household appliances and other non-hazardous materials. The quantity produced per person varies with the effectiveness of the material recovery scheme in place and with the affluence of the neighbourhood from which it is collected.

### Methodology

To estimate emissions from the incineration of municipal wastes the following methodology has

been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of waste combusted (t)

EF – emission factors (kg/t)

### Activity data

Amount on municipal waste incinerated has been obtained from Environmental Agency of the Republic of Slovenia. The data are available from the year 2002 only.

**Table 6.3.1 Amount of waste incinerated**

Year	Amount of waste (t)
2002	260
2003	235
2004	126
2005	294
2006	349
2007	686
2008	566
2009	649
2010	53
2011	260
2012	232
2013	141
2014	38
2015	53
2016	72
2017	135

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 6.3.2 Emission factors for municipal waste incineration and references**

Pollutant	Value	Unit	References
NO <sub>x</sub>	1071	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
SO <sub>x</sub>	87	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
CO	41	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
NM VOC	5,9	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1

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NH <sub>3</sub>	3	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
PM <sub>2.5</sub>	3	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
PM <sub>10</sub>	3	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
TSP	3	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
BC	0,105	g/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Cd	4,6	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Hg	18,8	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Pb	58	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
As	6,2	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Cr	16,4	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Cu	13,7	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Ni	21,6	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Se	11,7	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Zn	24,5	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Dioxins/ Furans	52,5	ng/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Benzo(a)pyrene	0,0084	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Benzo(b)fluoranthene	0,0179	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Benzo(k)fluoranthene	0,0095	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
Indeno(1,2,3-cd)pyrene	0,0116	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
HCB	0,0452	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1
PCB	3,4	ng/t	Emission Inventory Guidebook, 2016, 5.C.1.a Municipal waste incineration, pg 9, Table 3-1

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

Emissions from municipal waste incineration are extremely low for all pollutants. Contribution to total national emissions for all pollutants is below 0,001 %.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se, Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2002-2017.

### Future Improvements

No improvements are planned for next submission.

## 6.4 Hazardous waste incineration

NFR Code 5C1bii

### Introduction

This sector comprises the atmospheric emissions from the incineration of hazardous wastes. The composition of hazardous waste varies considerably. It includes any unwanted hazardous/chemical waste such as acids and alkalis, halogenated and other potentially-toxic compounds, fuels, oils and greases, used filter materials...

### Methodology

To estimate emissions from the incineration of hazardous wastes the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of waste combusted (t)

EF – emission factors (kg/t)

### Activity data

Amount on hazardous waste incinerated has been obtained from Environmental Agency of the Republic of Slovenia. The data are available for individual plant from yearly reports for the period 1990 - 2017.

**Table 6.4.1 Amount of waste incinerated**

Year	Amount of waste (t)	Year	Amount of waste (t)
1990	815	2004	1366
1991	815	2005	1325
1992	815	2006	1616
1993	815	2007	1987
1994	456	2008	2091
1995	268	2009	2585
1996	389	2010	2836
1997	73	2011	2860
1998	335	2012	2994
1999	1031	2013	6883
2000	1261	2014	8235
2001	1190	2015	11110
2002	946	2016	8993
2003	1382	2017	10906

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air



Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 6.4.2 Emission factors for hazardous waste incineration and references**

Pollutant	Value	Unit	References
NO <sub>x</sub>	0,87	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
SO <sub>x</sub>	0,047	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
CO	0,07	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
NM VOC	7,4	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
PM <sub>2.5</sub>	0,004	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
PM <sub>10</sub>	0,007	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
TSP	0,01	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
BC	0,00014	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
Cd	0,1	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
Hg	0,056	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
Pb	1,3	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
As	0,016	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
Ni	0,14	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
Dioxins/ Furans	1	µg I-TEQ/t	Plant specific
Total 4 PAHs	0,02	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1
HCB	0,002	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.i, 5.C.1.b.ii, 5.C.1.b.iv Industrial waste incineration, pg 10, Table 3-1

There is no information whether emission factors of particulate matter include or exclude condensable component.

## Emissions

Hazardous waste incinerators are not significant source of emissions. However, they are likely to be more significant emitters of dioxins, cadmium and mercury than many other sources. This depends on the type of waste, the combustion efficiency and the degree of abatement. Contribution of HCB emissions to total national emissions is about 4 %, for other pollutants is below 0,5 %. Only incineration of waste without energy recovery is included in the NFR sector 5C.

## Recalculations

Emissions of As and Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

## Category-specific QA/QC and verification

According to general 2017 in-depth EU NECD review 2017 recommendation EMEP/EEA Air

Pollutant Emission Inventory Guidebook, 2016 was used for emission calculations. Activity data was checked as well. Only incineration of waste without energy recovery is included in the NFR sector 5C. Incineration of waste with energy recovery is included in NFR sector 1A1a Public electricity and heat production as described in the IIR 2019 in the Chapter 3.1.1.

### Future Improvements

No improvements are planned for next submission.

## 6.5 Clinical waste incineration

NFR Code 5C1biii

### Introduction

This sector comprises the atmospheric emissions from the incineration of hospital wastes. Hospital waste includes human anatomic remains and organ parts, waste contaminated with bacteria, viruses and fungi, and larger quantities of blood.

### Methodology

To estimate emissions from the incineration of hospital wastes the following methodology has been adopted for individual pollutant:

$$E = m \times EF$$

E – emission (kg)

m – amount of waste combusted (t)

EF – emission factors (kg/t)

### Activity data

Amount on clinical waste incinerated has been obtained from Environmental Agency of the Republic of Slovenia. The data are available for individual plant from yearly reports for the period 1994 - 2017. There is no data available before that period.

**Table 6.5.1 Amount of waste incinerated**

Year	Amount of waste (t)	Year	Amount of waste (t)
1994	132	2006	108
1995	0	2007	160
1996	0	2008	148
1997	214	2009	193
1998	205	2010	671
1999	85	2011	660

<b>2000</b>	109	<b>2012</b>	578
<b>2001</b>	280	<b>2013</b>	524
<b>2002</b>	441	<b>2014</b>	267
<b>2003</b>	534	<b>2015</b>	195
<b>2004</b>	138	<b>2016</b>	299
<b>2005</b>	113	<b>2017</b>	245

## Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 6.5.2 Emission factors for clinical waste incineration and references**

Pollutant	Value	Unit	References
<b>NO<sub>x</sub></b>	2,3	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>SO<sub>x</sub></b>	0,54	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>CO</b>	0,19	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>NM VOC</b>	0,7	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>TSP</b>	17	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>BC</b>	0,391	kg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Cd</b>	8	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Hg</b>	43	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Pb</b>	62	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>As</b>	0,2	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Cr</b>	2	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Cu</b>	98	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Ni</b>	2	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Dioxins/ Furans</b>	1	µg I-TEQ/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>Total 4 PAHs</b>	0,04	mg/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>HCB</b>	0,1	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1
<b>PCB</b>	0,02	g/t	Emission Inventory Guidebook, 2016, 5.C.1.b.iii Clinical waste incineration, pg 8, Table 3-1

There is no information whether emission factors of particulate matter include or exclude condensable component.

## Emissions

The most significant pollutants from waste incineration process are heavy metals. A variety of

organic compounds, including dioxin, furans, chlorobenzenes, chloroethylenes and polycyclic aromatic hydrocarbons are also present in hospital waste or can be formed during the combustion and post-combination processes. Organics in the flue gas can exist in the vapour phase or can be condensed or absorbed on fine particulate. The relative proportion of emissions contributed by hospital waste incineration varies among pollutants. Emissions of Hg contribute about 6 % and HCB 5 %. Contributions of other pollutants are below 0,5 %.

### Recalculations

Emissions of As, Cr, Cu and Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1994-2017.

### Future Improvements

No improvements are planned for next submission.

## 6.6 Cremation

NFR Code 5C1bv

### Introduction

This sector comprises the atmospheric emissions from the incineration of human bodies in a crematorium. Incineration of animal carcass is not included.

### Methodology

To estimate emissions from cremation the following methodology has been adopted for individual pollutant:

$$E = N \times EF$$

E – emission (kg)

N – number of human bodies cremated

EF – emission factor (kg/body)

### Activity data

Activity data used for emission calculation is a number of cremations per year. The data on human bodies cremated have been obtained from two crematories operating in Slovenia. Share of cremations has been growing steadily and represents almost 80 % of deceased in Slovenia.

**Table 6.6.1 Number of cremations per year**

Year	Number of cremations	Year	Number of cremations
1990	5600	2004	12025

<b>1991</b>	5700	<b>2005</b>	12688
<b>1992</b>	5800	<b>2006</b>	12476
<b>1993</b>	5942	<b>2007</b>	13132
<b>1994</b>	6003	<b>2008</b>	13720
<b>1995</b>	6599	<b>2009</b>	14343
<b>1996</b>	6889	<b>2010</b>	14567
<b>1997</b>	7595	<b>2011</b>	14792
<b>1998</b>	8337	<b>2012</b>	15609
<b>1999</b>	9175	<b>2013</b>	15944
<b>2000</b>	9572	<b>2014</b>	15671
<b>2001</b>	9917	<b>2015</b>	16592
<b>2002</b>	10665	<b>2016</b>	16241
<b>2003</b>	11843	<b>2017</b>	17001

## Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used.

**Table 6.6.2 Emission factors for cremation and references**

Pollutant	Value	Unit	References
<b>NO<sub>x</sub></b>	0,825	kg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>SO<sub>x</sub></b>	0,113	kg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>CO</b>	0,140	kg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>NM VOC</b>	0,013	kg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>TSP</b>	38,56	g/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>PM<sub>10</sub></b>	34,7	g/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>PM<sub>2.5</sub></b>	34,7	g/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Cd</b>	5,03	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Hg</b>	1,49	g/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Pb</b>	30,03	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>As</b>	13,61	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Cr</b>	13,56	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Cu</b>	12,43	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Ni</b>	17,33	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Se</b>	19,78	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Zn</b>	160,12	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Dioxins/ Furans</b>	0,027	µg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1

<b>Benzo(a)pyrene</b>	13,20	µg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Benzo(b)fluoranthene</b>	7,21	µg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Benzo(k)fluoranthene</b>	6,44	µg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>Indeno(1,2,3-cd)pyrene</b>	6,99	µg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>HCB</b>	0,15	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1
<b>PCB</b>	0,41	mg/body	Emission Inventory Guidebook, 2016, 5.C.1.b.v Cremation, cremation of human bodies, pg 9, Table 3-1

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

The contribution of Hg emissions from cremation to the total national emissions is significant (15 %). Other pollutants are of less importance. They contributed less than 0,1 % to national totals. Although the number of cremations has grown considerably in recent years, emissions still do not affect significantly on the total national inventory.

### Recalculations

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

### Future Improvements

No improvements are planned for next submission.

## 6.7 Wastewater handling

Sectors covered in this chapter are:

NFR Codes:

5D1 Domestic wastewater handling

5D2 Industrial wastewater handling

### Introduction

This sector covers emissions from domestic and industrial waste water handling. Activities considered within this sector are biological treatment plants and latrines (storage tanks of human excreta, located under naturally ventilated wooden shelters).

### Methodology

To estimate emissions of NH<sub>3</sub> from latrines (domestic waste water handling) the following

methodology has been adopted:

$$E = N \times EF$$

E – emission (kg)

N – number of persons using latrines

EF – emission factor (kg/person/year)

To estimate emissions of NMVOC from industrial waste water treatment the following methodology has been adopted:

$$E = q \times EF$$

E – emission (mg)

q – quantity of waste water (m<sup>3</sup>)

EF – emission factor (mg/m<sup>3</sup> waste water)

### Activity data

For calculation of NH<sub>3</sub> emissions from latrines the relevant activity data is a number of inhabitants who use latrines. It is assumed that tenants of country houses with no water-flushed toilet have to use latrines outside the house. In 2017, about 0,1 % of Slovene population were not connected to any way of waste water treatment. Data on inhabitants included into various types of domestic wastewater treatment were obtained from Statistical Office of the Republic of Slovenia and the database on municipal wastewater treatment plants collected by the Slovenian Environment Agency. Number of inhabitants who use latrines is presented in Table 6.7.1.

**Table 6.7.1 Number of inhabitants who use latrines**

Year	Number of inhabitants	Year	Number of inhabitants	Year	Number of inhabitants	Year	Number of inhabitants
1990	442553	1997	330596	2004	119855	2011	14388
1991	427672	1998	310159	2005	80134	2012	12353
1992	408996	1999	305732	2006	60311	2013	10305
1993	390473	2000	294223	2007	40517	2014	8251
1994	376694	2001	284307	2008	20324	2015	6193
1995	363635	2002	271466	2009	18423	2016	4132
1996	346510	2003	259018	2010	16402	2017	4134

For calculation of NMVOC emissions from industrial waste water handling, the relevant activity data is the amount of industrial wastewater output. Data on amount of industrial waste water for the period 2004-2017 were obtained from database of monitoring industrial effluents collected by the Slovenian Environment Agency. For the period 1990 - 2005 values of quantity of waste water were estimated as described in National Inventory Report 2012, chapter Industrial waste water, pg 252-256. Wastewater output with regard to various industries is presented in Table 6.7.2.

Table 6.7.2 Wastewater output with regard to various industries

Year	Production of pulp and paper	Production of leather	Production of soft drinks and alcohol beverage	Production of food	Production of milk	Production of meat	Production of pharmaceutical products
	Wastewater output (m <sup>3</sup> )						
1990	17785835	909674	1993106	378570	1054778	1070278	
1991	15813639	778661	1897174	369069	1034204	1059647	
1992	13167759	736567	1773698	245566	921828	764296	
1993	12056736	686178	1812219	272168	767155	650592	
1994	13879156	678212	1906083	296905	835621	634050	
1995	15431625	459865	1879191	304715	911369	574572	
1996	14369458	529332	1881993	300437	885387	662932	
1997	16266638	496348	1941510	282961	926754	663706	
1998	18163843	463364	2001042	265483	968119	664480	
1999	20061023	430379	2060559	248007	1009486	665255	
2000	21397736	397395	2120086	230529	1050850	666029	
2001	22734450	364411	2179603	213054	1092218	666803	
2002	24071163	331427	2239130	195578	1133582	667578	
2003	25407851	298442	2298652	178100	1174950	668352	
2004	27672000	274700	1970685	136139	1133979	662367	1577989
2005	26947000	233185	1362038	178404	1230059	1420996	1368549
2006	21112000	238400	2074000	164120	986677	1143262	1544907
2007	12231000	281863	1771724	85040	984528	1393753	1487780
2008	16508000	228651	1572889	191920	981910	1334951	1523185
2009	15881919	11617	1533764	223853	901292	1162973	1765726
2010	13596494	9224	1737723	167710	865144	1268351	1633612
2011	12514742	22597	1785722	213732	871805	1161579	1560375
2012	12773572	39893	1543121	297757	820968	1119638	1465488
2013	10408933	44994	1458113	343151	835151	1074228	1528190
2014	11206175	47428	1268376	320628	838646	1144594	1578317
2015	11456759	40083	1166600	301864	750391	1307631	1684019
2016	11491537	35961	1058938	232644	805551	1724137	1747853
2017	11387032	45468	1031081	246433	854688	1727256	1783843

### Emission factors

A default emission factors for NH<sub>3</sub> and NMVOC were used for emission calculation. Emission factors were taken from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016.

Table 6.7.3 Emission factors for latrines and waste water handling

Pollutant		Value	Unit	References
NH <sub>3</sub>	Latrines	1,6	kg/person/year	Emission Inventory Guidebook, 2016, 5D Waste water handling, Table 3-2, pg 8
NMVOC	Waste water treatment in industry	15	mg/m <sup>3</sup> waste water	Emission Inventory Guidebook, 2016, 5D Waste water handling, Table 3-3, pg 9

### Emissions

Latrines are generally only a minor source of NH<sub>3</sub> emissions. The contribution of this activity to



the total ammonia emissions in the year 2017 is only 0,04 %. Drop of emissions in 2004 was due to wider inclusion of Slovene population into public sewage system in the last decade. More precise data are available for that period as well (Figure 6.7.1).

Biological treatment plants are only of minor importance for emissions into air, and the most important of these emissions are greenhouse gases CH<sub>4</sub>. Contribution of air pollutants to the total emissions is insignificant (0,001 %). Only very small quantities of NMVOC are emitted (Figure 6.7.2).

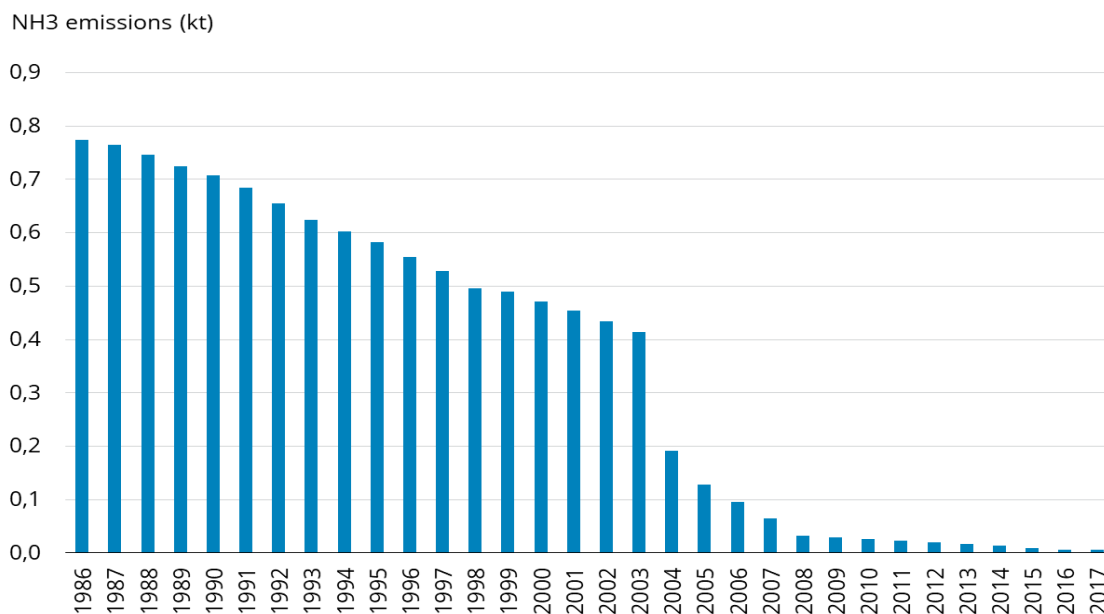


Figure 6.7.1 NH<sub>3</sub> emissions from latrines

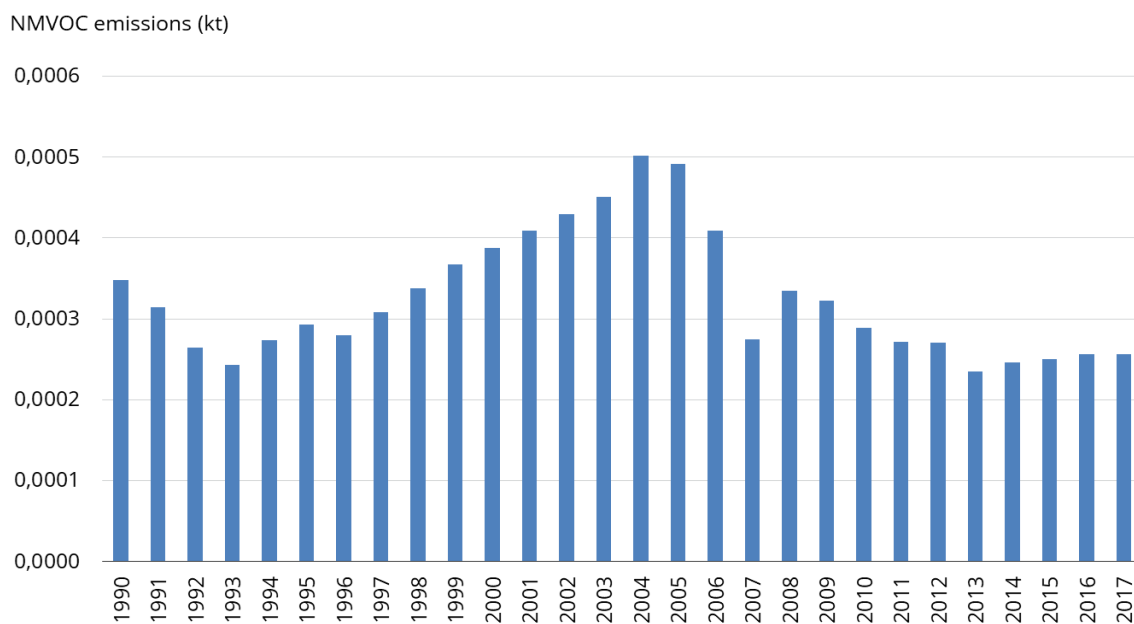


Figure 6.7.2 NMVOC emissions from industrial waste water treatment

## Recalculations

No recalculations have been performed since last submission.

## Future Improvements

No improvement is planned for this category.

## 6.8 Other waste

NFR Code 5E

### Introduction

This sector comprises emissions from car, house and industrial building fires. A limited amount of sludge was spread on the agriculture land and corresponding emissions have been included in the agriculture sector in category 3Da2b. There is no other evidence of sludge spreading in Slovenia.

### Methodology

To estimate emissions from fires the following methodology has been adopted for individual pollutant:

$$E = N \times EF$$

E – emission (kg)

N – number of fires

EF – emission factor (kg/fire)

### Activity data

Activity data used for emission calculation is a number of fires per year. Activity data for the period 2005-2017 has been provided by Administration for Civil Protection and Disaster Relief of the Republic of Slovenia. Data for the period 1990-2004 was estimated. Value of 2005 was used for emission calculation for the period 1990-2004.

**Table 6.8.1 Number of car, house and building fires per year**

Year	Number of car fires	Year	Number of house fires	Year	Number of industrial buildings fires
1990-2004	508	1990-2004	2040	1990-2004	25
2005	508	2005	2040	2005	25
2006	566	2006	2142	2006	3
2007	544	2007	2136	2007	9

<b>2008</b>	552	<b>2008</b>	2042	<b>2008</b>	8
<b>2009</b>	456	<b>2009</b>	2035	<b>2009</b>	15
<b>2010</b>	394	<b>2010</b>	1702	<b>2010</b>	125
<b>2011</b>	412	<b>2011</b>	1941	<b>2011</b>	207
<b>2012</b>	371	<b>2012</b>	1918	<b>2012</b>	169
<b>2013</b>	361	<b>2013</b>	1821	<b>2013</b>	164
<b>2014</b>	370	<b>2014</b>	1731	<b>2014</b>	159
<b>2015</b>	368	<b>2015</b>	1882	<b>2015</b>	151
<b>2016</b>	368	<b>2016</b>	1972	<b>2016</b>	162
<b>2017</b>	441	<b>2017</b>	2234	<b>2017</b>	184

### Emission factors

In calculating emissions of individual gases, following emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 have been used:

- for car fire: Table 3-2, pg 6, for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, dioxins/furans,
- for house fire: Table 3-4, pg 7, for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, Pb, Cd, Hg, As, Cr, Cu, dioxins/furans,
- for industrial building fire: Table 3-6, pg 8, for PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, Pb, Cd, Hg, As, Cr, Cu, dioxins/furans.

**Table 6.8.2 Emission factors for fires**

Car fires	Pollutant	Value	Unit	References
<b>Car fires</b>	<b>TSP</b>	2,3	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 6, Table 3-2
	<b>PM<sub>10</sub></b>	2,3	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 6, Table 3-2
	<b>PM<sub>2.5</sub></b>	2,3	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 6, Table 3-2
	<b>Dioxins/ Furans</b>	0,048	mg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 6, Table 3-2
<b>House fires</b>	<b>TSP</b>	61,62	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>PM<sub>10</sub></b>	61,62	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>PM<sub>2.5</sub></b>	61,62	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>Dioxins/ Furans</b>	0,62	mg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>Cd</b>	0,36	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>Hg</b>	0,36	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>Pb</b>	0,18	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>As</b>	0,58	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
	<b>Cr</b>	0,55	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4
<b>Cu</b>	1,28	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 7, Table 3-4	
<b>Industrial building fires</b>	<b>TSP</b>	27,23	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6

<b>PM<sub>10</sub></b>	27,23	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>PM<sub>2.5</sub></b>	27,23	kg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Dioxins/ Furans</b>	0,27	mg/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Cd</b>	0,16	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Hg</b>	0,16	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Pb</b>	0,08	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>As</b>	0,25	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Cr</b>	0,24	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6
<b>Cu</b>	0,57	g/fire	Emission Inventory Guidebook, 2016, 5E Other waste, pg 8, Table 3-6

There is no information whether emission factors of particulate matter include or exclude condensable component.

### Emissions

The contribution of emissions from fires to total national emissions is about 8 % for dioxins/ furans and 1 % for particulate matter. Contributions of heavy metals are less than 0,5 %. Emissions from this NFR sector were included into national inventory for the first time according to NECD 2017 review TERT recommendation.

### Recalculations

Emissions of As, Cr and Cu have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Emissions of Hg and Cd were recalculated for the period 1990-2016 due to error in emission calculation.

### Future Improvements

No improvements are planned for next submission.

**Biological treatment of waste - Anaerobic digestion at biogas facilities: NFR Code 5B2**  
**Industrial waste incineration: NFR Code 5C1bi**  
**Sewage sludge incineration: NFR Code NFR 5C1bi**  
**Other waste incineration (please specify in the IIR): NFR Code 5C1bvi**  
**Open burning of waste: NFR Code 5C2**  
**Other wastewater handling: NFR Code 5D3**

Notation Key "NO" (not occurring) were used for these sectors, since they are not sources of any additional emissions in Slovenia. No emissions occur in these sectors.

## 7 RECALCULATIONS AND IMPROVEMENTS

In general, considerable work has been carried out in the last few years to improve the inventory. New investigations and research carried out in Slovenia and abroad were, as far as possible, included as the basis for the emission estimates and included as data in the inventory databases. Furthermore, the updates of the EMEP/EEA air pollutant emission inventory guidebook and the work in the Task Force on Emission Inventories and Projections and its expert panels are followed closely in order to be able to incorporate the best scientific information as the basis for the inventories. Further important references in this regard are the 2006 IPCC Guidelines for National Greenhouse Gas Inventories. Implementation of new results in inventories is made in a way so that improvements better reflect Slovenia conditions and circumstances. In improving the inventories, care is taken to consider implementation of improvements for the whole time-series of inventories, to promote consistency. Such efforts lead to recalculation of previously submitted inventories.

In the last two years IIR was improved with better transparency of emission factors and activity data used and methodology applied. Our main goal was to calculate emissions according to revised guidelines for reporting emissions and projections data under the Convention LRTAP (ECE/EB.AIR/122/Add.1, decisions 2013/3 and 2013/4) and ensure completeness of the inventory. We focused great attention on introduction of new sources. We made a thorough examination of all emission factors used. We also pay special attention on notation keys used. NFR tables were corrected and filled with appropriate notation keys.

In June 2017 and 2018 our national inventory was subjected to in-depth EU NECD review. We improved our inventory with most of TERT expert review team recommendations. We applied the methodology and emission factors from new EMEP/EEA Emission Inventory Guidebook, 2016 for all sectors. Recalculation of emissions from all sectors were performed due to use of new guidebook and in-depth EU NECD review recommendations. A huge effort was put to check and implement all changes in emission factors and methodologies for all sectors. In Annex 3 to the IIR 2019 implementation of recommendations is included.

The main improvement in 2019 submission was introduction of additional heavy metals (As, Cr, Cu, Ni, Se, Zn) into national inventory. Emissions of additional heavy metals were estimated for all NFR sectors for the whole period 1990-2017.

Information on condensable component of particulate matter was introduced into IIR 2019. A table summarising whether PM<sub>10</sub> and PM<sub>2.5</sub> emission factors for each source sector include or exclude the condensable component and references for their emission factors are presented in Annex 2 to the IIR 2019. Indication in the methodology sections of IIR 2019 is included as well.

We are planning to use COPERT 5 and estimate uncertainty in next two years.

### 7.1 Recalculations

Recalculations in following sectors have been done since last submission to improve inventory:

#### Energy

##### Public electricity and heat production (1A1a)

According to 2018 in-depth EU NECD review imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory

Guidebook, 2016 were used for emissions calculation. Recalculation of emissions were therefore performed. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb and NMVOC were recalculated for the period 1990-2016. Recalculation of PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC, SO<sub>x</sub> and NO<sub>x</sub> emissions were performed for the period 2009-2016.

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

#### Petroleum refining (1A1b)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2003.

#### Manufacture of solid fuels and other energy industries (1A1c)

According to 2018 in-depth EU NECD review imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 were used for emissions calculation. Recalculation of emissions were therefore performed. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb, NMVOC and NO<sub>x</sub> were recalculated for the period 1995-2001. Recalculation of PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC emissions were performed for 2000 and 2001.

Due to new activity data on natural gas obtained for 2016 recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene and indeno(1,2,3-cd)pyrene was performed for the year 2016.

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

#### Manufacturing Industries and Construction (1A2)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Due to new activity data on natural gas obtained for 2016 in Other manufacturing industries and construction recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene and indeno(1,2,3-cd)pyrene was performed for the year 2016.

Activity data on anthracite was obtained for 1996 in Other manufacturing industries and construction sub sector. Recalculation of SO<sub>x</sub>, NO<sub>x</sub>, NMVOC, CO, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, BC, Pb, Cd, Hg, dioxins/furans, benzo(a)pyrene, benzo(b)fluoranthene, benzo(k)fluoranthene indeno(1,2,3-cd)pyrene, HCB and PCB was performed for the year 1996.

#### Road transport (1A3b)

According to 2018 in-depth EU NECD review emissions of Pb and Cd from automobile tyre and brake wear have been estimated for the whole period 1990-2017. Copert 4 (version 11.4) have been used for estimation of non-exhaust emissions.

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017. Exhaust and non-exhaust emissions have been estimated.

Emission of SO<sub>x</sub> was recalculated for 2016 due to error in filling the NFR reporting table.

#### Railways (1A3c)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Imported sub-bituminous coal should be classified as hard coal. Corresponding emission factors from EMEP/EEA Air Pollutant Emission Inventory Guidebook, 2016 were used for emissions calculation. Emissions of benzo(a)pyrene, indeno(1,2,3-cd)pyrene, Hg, Cd, Pb and NMVOC were recalculated for the period 1990-2016. Recalculation of PM<sub>2.5</sub>, PM<sub>10</sub>, TSP, BC were performed for the period 2000-2016 and recalculation of SO<sub>x</sub> and NO<sub>x</sub> for the period 1980-2016.

International maritime navigation (1A3di(i))

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2006-2017.

Residential: Stationary (1A4bi)

Commercial/institutional: Stationary (1A4ai)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Data on wood consumption in 1A4bi Residential: Stationary for the period 1990-2016 has been improved and related emissions have been recalculated for that period.

Mobile Combustion in manufacturing industries and construction (1A2gvii)

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Agriculture/Forestry/Fishing: Off-road vehicles and other machinery (1A4cii)

Emissions of Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Pipeline transport (1A3ei)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2008-2017.

Fugitive emissions oil: Refining / storage (1B2aiv)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2001.

Venting and flaring (oil, gas, combined oil and gas) (1B2c)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

**Industrial processes and product use**

Glass production (2A3)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Titanium dioxide production (2B6)

According to 2018 in-depth EU NECD review recalculation of SO<sub>x</sub> emissions were performed for the period 1982-2016. Tier 3 plant specific data was used for estimation of SO<sub>x</sub> emissions.

Chemical industry: Other (2B10a)

Emissions of SO<sub>x</sub>, NMVOC and TSP have been recalculated for the years 2014 and 2015 due to mistake in calculations made in previous submission..

Iron and steel production (2C1)

Emissions of As, Cr, Cu, Ni and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Due to change in emission factor used for period 1988-1993 emissions of CO, NO<sub>x</sub> and SO<sub>x</sub> have been recalculated for the period 1988-1993, emissions of NMVOC for the period 1990-1993 and emissions of dioxins/furans, PAHs, PCB, Pb, Cd, Hg for the period 1990-1993.

Aluminium production (2C3)

Emissions of dioxins/furans and HCB have been calculated and included into the national



inventory for the first time. Emissions of dioxins/furans were introduced for the period 1990-2017, emissions of HCB for the period 1990-2001.

Lead production (2C5)

Emissions of As and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Zinc production (2C6)

Emissions of As and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Copper production (2C7a)

Emissions of As, Cr, Cu, Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Asphalt roofing (Code 2D3c)

Since 2009 NMVOC emissions have been recalculated using data from the Remis database.

Coating Application (2D3d)

For the period 2011-2016 NMVOC, emissions have been recalculated using data on amount of NMVOC from the tax database.

Chemical Products (2D3g)

Due to the obvious error in AD for 2001, NMVOC emissions for this year have been interpolated.

Other solvent and product use (2D3i and 2G)

Following the recommendation from the peer review NMVOC emissions from airplane de-icing have been included in the inventory for the period 1990-2017.

In addition, emissions of six heavy metals (As, Cr, Cu, Ni, Se, and Zn) for the period 1990-2017 have been included in the inventory for the first time.

Pulp and paper industry (2H1)

Since 2006 emissions of NMVOC have been recalculated using measurements.

Food and beverages industry (2H2)

NMVOC emissions for the whole period have been recalculated because emissions from spirits have been excluded by mistake in the previous submission.

## **Agriculture**

Inorganic N-fertilizers (3Da1)

Error in calculation of ammonia emissions from urea application was discovered in previous report (factor 0,015 was used instead of 0,155). The value was corrected. As a result, ammonia emissions for the entire reporting period increased.

On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied.

As a result the emissions of nitric oxide were reduced for the entire reporting period.

Animal manure applied to soils (3Da2a)

The ammonia emissions for 2015 and 2016 were corrected by taking into account the new information on the low emission manure spreading techniques which were supported by Rural development programme. As a result, the emissions of ammonia for the above mentioned years were reduced.

On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040

kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced for the entire reporting period.

#### Sewage sludge applied to soils (3Da2b)

The emission factor 0,81 kg of ammonia nitrogen per kg of total ammonia nitrogen applied by sewage sludge, which was used in previous report, was replaced by a factor 0,13 kg of ammonia nitrogen per kg of total ammonia nitrogen applied by sewage sludge (EMEP/EEA 2016). It resulted in a decrease of ammonia emissions. Recalculations for the entire period were done. On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced.

#### Urine and dung deposited by grazing animals (3Da3)

Recalculations for the entire period were done. On the initiative of reviewers, the emission factor, which was previously misinterpreted as 0,040 kg NO per kg of N, was replaced by a factor 0,040 kg NO<sub>2</sub> per kg of N applied. As a result, the emissions of nitric oxide were reduced.

#### Farm-level agricultural operations including storage, handling and transport of agricultural products (3Dc)

It was found that for 1990 the calculation procedures were not properly linked to data on the areas of arable land, temporary and permanent grasslands. As a result emissions from soil cultivation, harvesting, cleaning and drying of crops were not estimated. Error was eliminated. It did not affect the results for the period 1991-2016.

## **Waste**

#### Biological treatment of waste - Solid waste disposal on land (5A)

Recalculations of NMVOC, PM<sub>2.5</sub>, PM<sub>10</sub> and TSP were recalculated for the year 2016 due to new data on amount of landfilled waste obtained.

#### Municipal waste incineration (5C1a)

Emissions of As, Cr, Cu, Ni, Se, Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 2002-2017.

#### Hazardous waste incineration (5C1bii)

Emissions of As and Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

#### Clinical waste incineration (5C1biii)

Emissions of As, Cr, Cu and Ni have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1994-2017.

#### Cremation (5C1bv)

Emissions of As, Cr, Cu, Ni, Se and Zn have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

#### Other waste (5E)

Emissions of As, Cr and Cu have been calculated and included into the national inventory for the first time. Emissions were introduced for the whole period 1990-2017.

Emissions of Hg and Cd were recalculated for the period 1990-2016 due to error in emission calculation.

**Table 7.1.1 Changes due to recalculations of main pollutants emissions between 2019 and 2018 inventory submission for inventory year 2016**

Sector	Main Pollutants				Other
	NO <sub>x</sub> (as NO <sub>2</sub> )	NM VOC	SO <sub>x</sub> (as SO <sub>2</sub> )	NH <sub>3</sub>	CO
	kt	kt	kt	kt	kt
1A1 Energy industries	-0,00967	-0,00266	-0,24229	NE	-0,00424
1A2 Manufacturing industries and construction	0,00805	0,00250	0,00007	0,00000	0,00318
1A3 Transport	-0,00015	0,00000	0,00298	0,00000	0,00000
1A4 Small combustion and non-road mobile sources and machinery	0,00001	0,00003	0,00000	0,00000	0,00025
1A5 Other	0,00000	0,00000	0,00000	0,00000	0,00000
1B Fugitive emissions from fuels	0,00000	0,00000	0,00000	NA	0,00000
2A Mineral industry	NE	NE	NE	NE	NE
2B Chemical industry	0,00000	0,00000	-0,17494	NE	NE
2C Metal industry	0,00000	0,00000	0,00000	NE	0,00000
2D-2L Other solvent and product use	0,00083	-0,54946	0,00000	0,00192	0,02548
3B Manure management	0,00000	-0,05455	NA	0,00000	NA
3D-3I Crop production and agricultural soils	-1,22685	0,00000	NA	0,56713	NA
5A Biological treatment of waste - Solid waste disposal on land	NA	0,00000	NA	NE	NE
5B Biological treatment of waste - Composting	NE	NE	NE	0,00000	NE
5C Waste incineration	0,00000	0,00000	0,00000	0,00000	0,00000
5D Wastewater handling	NA	0,00000	NA	0,00000	NA
5E Other waste	NE	NE	NE	NE	NE

**Table 7.1.2 Changes due to recalculations of particulate matter emissions between 2019 and 2018 inventory submission for inventory year 2016**

Sector	Particulate Matter			
	PM <sub>2.5</sub>	PM <sub>10</sub>	TSP	BC
	kt	kt	kt	kt
1A1 Energy industries	-0,00010	-0,00010	-0,00010	0,00000
1A2 Manufacturing industries and construction	0,00009	0,00009	0,00009	0,00000
1A3 Transport	0,00000	0,00000	0,00000	0,00000
1A4 Small combustion and non-road mobile sources and machinery	0,00003	0,00003	0,00003	0,00000
1A5 Other	NE	NE	NE	NE
1B Fugitive emissions from fuels	0,00000	0,00000	0,00000	0,00000
2A Mineral industry	0,00000	0,00000	0,00000	0,00000
2B Chemical industry	0,00000	0,00000	0,00000	NA

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2C Metal industry	0,00000	0,00000	0,00000	0,00000
2D-2L Other solvent and product use	0,01218	0,01097	0,00641	0,00006
3B Manure management	0,00000	0,00000	0,00000	NA
3D-3I Crop production and agricultural soils	0,00000	0,00000	NA	NA
5A Biological treatment of waste - Solid waste disposal on land	0,00000	0,00000	0,00000	NA
5B Biological treatment of waste - Composting	NE	NE	NE	NE
5C Waste incineration	0,00000	0,00000	0,00000	0,00000
5D Wastewater handling	NE	NE	NE	NE
5E Other waste	0,00000	0,00000	0,00000	NE

**Table 7.1.3 Changes due to recalculations of heavy metals emissions between 2019 and 2018 inventory submission for inventory year 2016**

Sector	Priority Heavy Metals		
	Pb	Cd	Hg
	t	t	t
1A1 Energy industries	-0,04585	-0,00536	-0,00894
1A2 Manufacturing industries and construction	0,00001	0,00000	0,00006
1A3 Transport	-0,00003	0,00000	0,00000
1A4 Small combustion and non-road mobile sources and machinery	0,00000	0,00000	0,00000
1A5 Other	NE	NE	NE
1B Fugitive emissions from fuels	0,00000	0,00000	0,00000
2A Mineral industry	0,00000	0,00000	0,00000
2B Chemical industry	NE	NE	NE
2C Metal industry	0,00000	0,00000	0,00000
2D-2L Other solvent and product use	0,34665	0,00250	0,00003
3B Manure management	NA	NA	NA
3D-3I Crop production and agricultural soils	NA	NA	NA
5A Biological treatment of waste - Solid waste disposal on land	NA	NA	NE
5B Biological treatment of waste - Composting	NA	NA	NA
5C Waste incineration	0,00000	0,00000	0,00000
5D Wastewater handling	NE	NE	NE
5E Other waste	0,00000	0,00001	0,00001

**Table 7.1.4 Changes due to recalculations of POPs emissions between 2019 and 2018 inventory submission for inventory year 2016**

Sector	POPs							
	PCDD/ PCDF (dioxins/ furans)	PAHs					HCB	PCBs
		benzo(a) pyrene	benzo(b) fluoranthene	benzo(k) fluoranthene	Indeno (1,2,3-cd) pyrene	Total 1-4		
	g I-TEQ	t	t	t	t	t	kg	kg
1A1 Energy industries	-0,00005	0,00000	0,00000	0,00000	-0,00001	-0,00001	0,00000	0,00000
1A2 Manufacturing industries and construction	0,00006	0,00008	0,00032	0,00012	0,00012	0,00063	0,00000	0,00000
1A3 Transport	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000
1A4 Small combustion and non-road mobile sources and machinery	0,00004	0,00001	0,00000	0,00000	0,00000	0,00002	0,00000	0,00000
1A5 Other	NE	NE	NE	NE	NE	NE	NA	NA
1B Fugitive emissions from fuels	NA	NA	NA	NA	NA	NA	NA	NA
2A Mineral industry	NA	NA	NA	NA	NA	NA	NA	NA
2B Chemical industry	NA	NA	NA	NA	NA	NA	NA	NA
2C Metal industry	0,05906	0,00000	0,00000	0,00000	0,00000	0,00000	NE	0,00000
2D-2L Other solvent and product use	0,00005	0,00005	0,00002	0,00002	0,00002	0,00011	NE	0,00000
3B Manure management	NA	NA	NA	NA	NA	NA	NA	NA
3D-3I Crop production and agricultural soils	NA	NA	NA	NA	NA	NA	NA	NA
5A Biological treatment of waste - Solid waste disposal on land	NA	NA	NA	NA	NA	NA	NA	NA
5B Biological treatment of waste - Composting	NA	NA	NA	NA	NA	NA	NA	NA
5C Waste incineration	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000	0,00000
5D Wastewater handling	NA	NA	NA	NA	NA	NA	NA	NA
5E Other waste	0,00000	NE	NE	NE	NE	NE	NE	NE

## 7.2 Planned improvements

### Road transport (1A3b)

We are planning to use new COPERT 5 model for emission calculation from road transport in next two years.

### Domestic solvent use including fungicides (2D3a)

To calculate NMVOC emissions from domestic use of solvents with Tier 2 approach at least the following data on product or solvents use is needed:

- Cosmetics and toiletries: hair sprays, toilet waters, after shaves, perfumes, face cares, personal deodorants and antiperspirants, body care products
- Household products: polishes and creams, soaps
- Car care products: antifreeze

National data on the use of these products is not available.

In the 2016 guidelines, it is proposed that the Tier 2 method has to be used in the EU countries for the assessment, as there are data on the use of solvents in the solvent manufacturer database. In the document (ESIG, 2015), which is indicated in the instructions as a source for

data on the use of solvents of solvent-containing products, Slovenia and Austria are taken into account together, therefore data for Slovenia is not available. Because GDP per capita in Slovenia is much smaller than in Austria (in 2017 23,596\$ and 47,221\$ per capita, respectively), comparison with Austria is not suitable.

In addition, the ESIG inventory suggests an emissions factor of 1.2 kg/capita to be used for the EU-27. This EF is used in our calculation of NMVOC emissions.

We will try to estimate activity data, using per capita data from similar country (or group of countries) for the next submission.

#### Coating Application (2D3d)

For the next submission, we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

The next step of improvement for this category would be a split of decorative coating application between domestic use and paint use in construction and buildings. At that moment, we have no data and no reliable methodology to perform such disaggregation. As this improvement would have no effect on the total emissions, it is not planned for the near future.

#### Degreasing (2D3e)

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

#### Chemical Products (2D3g)

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

Following the TERT recommendation, we will obtain data on NMVOC (total organic o-toluidine) emissions from Remis database since 2009 for relevant plastics production processes and assess the implementation of this methodology for the next submission.

#### Printing (2D3h)

For the next submission we are planning to use improved data from the HOS database. For more information, please see description on the future improvements in the Chapter 4.4.10 Other solvent and product use.

#### Other solvent and product use (2D3i and 2G)

For category 2D3i Other Solvent Use and pollutant NMVOC for year 2006 the TERT noted that there was a sharp increase of NMVOC emissions in 2006 by 2.8 times compared to the year 2005. After investigation we have found out, that HOS database was not completed for production of glues for 2005, 2006 and 2008. For this reason, we have rechecked all data in the HOS database for all years and all categories and some other missing data have been found. Now, the database is complete, but was not complete at the time of inventory submission on 15 February. Therefore, the recalculation of emissions from HOS database in the year 2005, 2006, and 2008 will be done for the next submission. This recalculation will also affect emissions for the period 1990-2004, because in some cases when the data for the years before 2005 are not available, emissions in the in 2005 was used to extrapolate emissions back to 1990.

## 8 PROJECTIONS

Projections reported in 2019 are the same as have been reported in 2017. New projection are being prepared and will be available at the end of 2019.

### 8.1 Bases for the preparation of projections

For the preparation of emission projections from energy sources, the results of long-term energy balances prepared for the period 2012 - 2030 for the Ministry of the Economy have been used as a basis. Within the framework of balances, several scenarios were used, composed of three scenarios of economic development, two strategies for the implementation of measures (in the field of energy efficiency and renewable energy sources), two scenarios of transport development and one scenario for the development of the electrical-energy sector. The with measures scenario was composed of moderate scenario of economic growth (scenario +), reference scenario of the implementation of measures and a balanced scenario of energy supply. The base year for the preparation of long-term balances was 2011, but for some sectors models were calibrated also for other years e.g. for transport for years 2008-2012, for households for years 2009-2012, etc. For the 2016 emissions projections activity projections from 2012 have been slightly corrected based on the recent activity developments but long-term projections have remained the same.

In the course of the energy projections calculation of the shares of different technologies have been determined e.g. wood combustion in households in stoves, automatic boilers, old manual boilers, new manual boilers, vehicle fleet for cars, trucks, buses, etc.. This has important implications for the emissions calculation.

Activity data in industrial processes have been calculated on the basis of the historical development and projections of economic activity (production index) that was also used for energy projections.

For the emissions from solvents use in industry production indexes for different branches were used, while for other sources population has been used as a proxy.

The basis for the preparation of projections in agriculture is the agriculture development strategy, while for the waste sector operational programme on urban waste management.

### 8.2 Methodology of activity projections

#### Energy

In order to prepare emissions projections for the energy sector as defined by the guidelines, a system of models was used in which the main tool is a reference energy ecological model called REES-SLO2, made in the MESAP environment. The technology-orientated "Reference Energy-Ecological Model for Slovenia (REES-SLO2)" was developed in the MESAP environment in the form of a linear network model for processes and connections (a reference energy system), which enables consistent modelling of energy use based on the needs of energy services and energy supply according to the Integrated Resource Planning method. The tools, models and methodology were verified in a series of preliminary studies and have been used as a basis for many strategic documents for the development of the energy sector and the reduction of greenhouse gas and pollutants emissions in Slovenia. The set of models includes the following submodels:

Firstly, with the help of a model for the assessment of the market penetration of energy saving and renewable energy technologies (PET SLO), the market shares of individual energy efficient and renewable technologies per final uses are calculated as a response to changing price signals, financial incentives and information campaigns. Minimum requirements regarding energy efficiency (of buildings, equipment, products) are taken into account in PET SLO model as boundary conditions determining available technologies. Efficient use of energy measures in

the energy-intensive fields of the processing industry are also modelled separately. Result of PET SLO model are the market shares of certain technologies per energy use.

REES-SLO2 (implemented in MESAP) calculates the envisaged final energy use balances and assesses the local production of electricity and heat based on the proportions of different technologies in the final use structure and connections with influential parameters (the levels of economic activity by sector, the number of households, etc.). The final use of electricity divided by sector, purpose, and production in local supply systems (in industrial, distribution, and private units) is transferred for processing by the program in order to analyse the load shape.

With the help of a model for the optimisation of electricity production in free market conditions, the total production and use of electricity and system prices are calculated, as well as the quantities to be provided by individual major producers. The calculation is based on the optimisation of all supply offers from producers considering prices on international markets, taking into account the technical limitations of individual facilities and the objectives regarding the stability of the system.

The proportions of electricity production in individual units calculated in Point 3 and related costs are transferred to the MESAP/REES-SLO2 model. Other balances are calculated for the whole planning period in the MESAP model: primary and final energy, emissions (CO<sub>2</sub>, CH<sub>4</sub>, N<sub>2</sub>O, SO<sub>2</sub>, NO<sub>x</sub>, NMVOC, TSP, PM<sub>10</sub>, PM<sub>2.5</sub>, NH<sub>3</sub>) and total costs.

### Transport

Two models were used for transport emissions. For the assessment of the final energy consumption, an energy model for transport was prepared. The basis for the calculation of energy use in transport was an estimation of the development of the transport activity. This was done on the basis of the results of study prepared by PNZ for the Ministry responsible for transport and has been used for the preparation of the transport strategy. Two scenarios have been prepared – continuation of the past trend meaning roads centric development and balanced transport development, with increase in public transport and rail transport. Additional model calculates the vehicle fleet structure for different types of vehicles on the basis of assumptions regarding vehicle life-cycle and the structure of newly-purchased vehicles. Boundary condition for the shares of different technologies in newly purchased cars was set on the basis of the EU Regulation 443/2009 setting emission performance standards for new passenger cars as part of the Community's integrated approach to reduce CO<sub>2</sub> emissions from light-duty vehicles. From the assumption of the shares of various types of transport and the technical characteristics of vehicle fleets, the model calculates the energy use. Transit transport has an important effect on the amount of fuel sold in Slovenia coupled with the market situation of fuel prices in Slovenia and neighbouring countries. That is why transit transport is modelled separately.

Transport emissions were determined using the emissions factors of the 2016 GAINS model of the IIASA institute.

### Industrial processes and product use

The projections of emissions in industrial processes were made on the basis of projection of an industrial production index, taking different emission factors for different activities into account. In the projection of the primary aluminium production, a study of the Talum Company, which is the only primary aluminium producer in Slovenia, was taken into account. For PM and NMVOC emissions the same methodology was used as is used for the inventory preparation, with the exception of the sources where measurements are used in the inventory. In the projections emissions have been calculated as a product of activity and emission factor, because this is the only way which allows assessment of the effect of different measures.

Emissions from solvent use have been estimated with the same methodology as is used for the inventory preparation, with the exception of emissions from paint application and domestic solvent use and sources where reported/measured emissions are used in the inventory. For these sources activity data have been multiplied with emission factors which have been determined based on the GAINS guidelines for inventory preparation numbers.



### Agriculture

Agriculture emission projections were carried out according to the methodology prescribed by the latest EMEP/ EEA guidebook. The EMEP/ EEA methodology anticipates agriculture emission projections based on statistical data on the physical volume of crop and animal production, taking into account specific procedures characteristic of particular countries or areas. Data on the extent of crop production and domestic animal production is treated separately, despite their interdependence. The model based on this methodology therefore does not enable optimisation at the level of the agriculture sector as a whole, but only with regard to separate segments. SORS statistical data and information obtained from experts in the agricultural sector were used for the assessment.

### Waste

Emissions from waste landfills have been estimated on the basis of the projections of the landfilled biodegradable waste. This was calculated from the amount of waste produced using shares of different waste manipulation. In the calculation reduction in the depositing of biodegradable waste is envisaged. The composition of the biologically degradable part was constant and calculated according to the results of screening analyses in Slovenia.

## **8.3 Uncertainty of projections**

The uncertainty of the projections arises from the uncertainty of the statistical data used as a basis for the projections (statistical data, emission factors), the models used for the projections, which present a simplified picture of the actual events, the uncertainty of the scenarios for the implementation of policies and measures since they change over time; furthermore, it is difficult to envisage the actual impact of measures, since they are influenced by many factors as well as the uncertainties of future economic, technological and social developments, including the uncertainty of energy prices, growth in the energy supply and demand, the behaviour of the main players in the energy market, etc.

The results of the emission projections in the energy sector are largely dependent on the realisation of the considered measures in the area of RES and EE, which will be dependent on the available budgetary funds and also staff in the ministries. The dynamics of the transition to natural gas in electricity production are largely dependent on future market movements and social problems as regards reducing the production of coal. Fugitive emission projections are also uncertain as a result of the uncertainty of the future development of the coal mining industry. Other sources of uncertainty include the scenarios for the future development of the gross domestic product, which has a strong impact on energy use and consequently on emissions from industry. Between 2020 and 2030, an average annual growth rate of 2.7 % is envisaged. For Slovenia, the transport sector represents the largest uncertainty in the preparation of projections. Projections of transport volume carry some uncertainty. The largest uncertainty with regard to projections in transport is represented by transit transport, which cannot be captured by models covering Slovenia only, since the transit flows originate elsewhere. Transit transport has significant influence on the sale of motor fuel in Slovenia coupled with fuel prices in Slovenia and the neighbouring countries, since vehicles in transit transport fill their tanks where fuel is the cheapest. The estimation of the share of fuels sold to foreign transit transport in the total fuel used in road transport in 2008 amounts to 30 % and in 2012 to 28 %. In 2015 share of foreign transit transport is estimated to 21 %.

Uncertainties in estimating emissions in agriculture arise in particular from the uncertainty regarding the fluctuation in the number of animals. The above-mentioned number changed considerably over recent years and for the future the policy of the Government of the Republic of Slovenia as regards increasing self-sufficiency in food was taken into account.

## 9 ABBREVIATIONS

AD	activity data
Al <sub>2</sub> O <sub>3</sub>	aluminium oxide
As	arsenic
BC	black carbon
BAT	best available techniques
C	confidential
CaO	calcium oxide
CaCO <sub>3</sub>	calcium carbonate
Cd	cadmium
CDR	Central Data Repository (of the EEA's Eionet Reportnet)
CEIP	Centre on Emission Inventories and Projections
CH <sub>4</sub>	methane
CLRTAP	(UNECE) Convention on Long-range Transboundary Air Pollution
CNG	compressed natural gas
CO	carbon monoxide
CO <sub>2</sub>	carbon dioxide
CORINAIR	COoRdination of INformation on AIR emissions
Cr	chromium
CRF	common reporting format (for greenhouse gases, UNFCCC)
CAS	Chemical Abstracts Service
COPERT	model and methodology for determination of road transport emission
CS	country specific
Cu	copper
D	default value
EC	European Commission
EEA	European Environment Agency
EF	emission factor
EIONET	European environmental information and observation network
EMEP	European Monitoring and Evaluation Programme
ETS	Emission Trading Scheme
EU	European Union
EURO	European emission standards define the acceptable limits for exhaust emissions of new vehicles sold in EU
EUROSTAT	Statistical Office of the European Communities
GHG	greenhouse gases
GB	EMEP/EEA Air Pollutant Emission Inventory Guidebook
FGD	device for the desulphurization of flue gases
Fe <sub>2</sub> O <sub>3</sub>	iron (III) oxide
HCB	hexachlorobenzene
HCE	hexachloroethane
HOS database	Slovenian database with plant specific emission values
Hg	mercury
HM(s)	heavy metal(s)
IE	included elsewhere
IEA	International Energy Agency
IED	Industrial Emissions Directive
IIR	Informative Inventory Report
IPCC	Intergovernmental Panel on Climate Change
IPPC	Integrated pollution prevention and control (EU Directive)
ISEE	Slovenian emission inventory information system
I-TEQ	international toxic equivalents

JQ	Joint Questioner, statistics data
KCA	key category analysis
LEG	annual energy statistics of the energy sector
LPG	liquefied petroleum gas
LRTAP	Long-range Transboundary Air Pollution
LTO	landing and take-off cycle, aviation
M	model
MgO	magnesium oxide
MSW	municipal solid waste
N	nitrogen
NCV	net caloric value
N <sub>2</sub> O	nitrous oxide
NA	not applicable
NE	not estimated
NECD	National Emission Ceilings Directive (2001/81/EC)
NFR	nomenclature for reporting (air pollutants, UNECE)
NH <sub>3</sub>	ammonia
Ni	nickel
NIR	National Inventory Report
NK	notation key
NMVOC(s)	non-methane volatile organic compound(s)
NO	not occurring
NO <sub>2</sub>	nitrogen dioxide
NO <sub>x</sub>	nitrogen oxides
NR	not relevant
O <sub>3</sub>	ozone
PAH(s)	polycyclic aromatic hydrocarbon(s)
Pb	lead
PCB(s)	polychlorinated biphenyl(s)
PCDD/F(s)	polychlorinated dibenzodioxin(s)/dibenzofuran(s)
PCDD	polychlorinated dibenzo-p-dioxins
PCDF	polychlorinated dibenzofurans
PCT	polychlorinated terphenyls
PM	particulate matter
PM <sub>10</sub>	coarse particulate matter (particles measuring 10 µm or less)
PM <sub>2.5</sub>	fine particulate matter (particles measuring 2.5 µm or less)
POP(s)	persistent organic pollutant(s)
PS	plant specific
QA	quality assurance
QC	quality control
REMIS database	Slovenian database with plant specific emission values
RS	Republic of Slovenia
SCA	Standard Classification of Activities
S	suphur
Se	selenium
SEA	Slovenian Environment Agency
SiO <sub>2</sub>	silicon dioxide
SNAP	Selected Nomenclature for reporting of Air Pollutants
SORS	Statistical Office of the Republic of Slovenia
SO <sub>2</sub>	sulphur dioxide
SO <sub>x</sub>	sulphur oxides
T	tier (method)
TERT	Technical Expert Review Team – 2017 NECD review
TAN	total ammonia nitrogen

TFEIP	Task Force on Emission Inventories and Projections
TSPs	total suspended particulates
UNECE	United Nations Economic Commission for Europe
UNFCCC	United Nations Framework Convention on Climate Change
VOC	volatile organic compound
Zn	zinc

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