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Atmospheric emissions of some POPs in Europe – a discussion of existing inventories and data needs

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20 Abstract

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22 Accurate and complete information on the emissions of persistent organic 23 pollutants (POPs) are essential for interpreting historical, current and future 24 contamination levels in remote areas. However, this information is also crucial 25 for decision makers aiming at further reduction of the environmental loading of 26 these substances on a regional scale. In this study, we identify and discuss 27 specific POP emission inventory features that are deemed essential to 28 understand, predict and control the behaviour of such substances on a 29 European scale. It is shown that the scientific value of official emission data is 30 limited (e.g. for deriving source-receptor relationships on a European scale), as there is insufficient information on spatial, temporal and compound 31 32 coverage. Likewise, we argue that non-official emission data (i.e. research-33 driven emission inventories), which are based on aggregated statistics, may 34 be of limited value for the identification of further emission control strategies. It 35 is thus argued that future emission inventories should be developed in a format that is suitable to serve both policy- and research oriented applications. 36 37 Further improvement of official emission data with respect to research-driven features seems to be the most sensible way to proceed. Finally, the empirical 38 39 basis of current emission inventories remains weak, and further research on emission identification and characterisation seem needed (a) to gain 40 confidence in predicted source-receptor relationship as well as (b) for the 41 42 development of sound control strategies.

44 **1. Introduction**

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Persistent Organic Pollutants (POPs) are chemicals that are recognised as persistent, bio-accumulative, toxic and susceptible to long-range atmospheric transport (PBT-LRT). POPs generally fall into two classes dependent on their origin; intentionally produced chemicals (typically organochlorinated pesticides and industrial chemicals) and unwanted by-products of combustion (polyaromatic hydrocarbons – PAHs, and dioxins - PCDD/Fs).

Elevated concentrations of several POPs have been observed in remote 52 environments far from global source regions, such as the Arctic (AMAP, 53 54 2004). Ottar (1981) hypothesised that several chlorinated hydrocarbons have 55 a potential for reversible atmospheric deposition, and that they therefore may 56 be subject to a systematic long-term transport from warmer to colder regions. 57 This theory was further elaborated by Wania and Mackay (1993) who offered more detailed explanations how the temperature-dependent partitioning of 58 59 persistent and low volatility compounds could lead to an enrichment in the 60 Arctic through global fractionation and cold condensation processes. Later, 61 several modelling studies have been undertaken to understand and predict source-receptor relationships for POPs on a European (e.g. VanJaarsveld et 62 al., 1997; Prevedouros et al., 2004) hemispherical (e.g. Malanichev et al., 63 64 2004; Hansen et al., 2004) and global scale (e.g. Wania and Su, 2004). 65 International agreements are now also coming into force to reduce further 66 environmental exposure of POPs on a regional and global scale. One of such international agreements is the Stockholm Convention on POPs (UNEP, 67 2001). The Stockholm Convention entered into force in May 2004 (151 68 69 signatories and 118 parties as of February 17, 2006). The Stockholm 70 Convention targets 12 POPs (the so-called "dirty dozen") for reduction and 71 eventual elimination, and sets up a system for evaluation of additional chemicals for consideration. Another international agreement is the 1979 72 Geneva Convention on long-range transboundary air pollution (LRTAP), which 73 74 has 49 parties (UN/ECE, 1979). The LRTAP convention has been extended 75 by the 1998 Aarhus protocol on Persistent Organic Pollutants (POPs), which 76 entered into force by the end of 2003 (25 ratifications as of January 13, 2006). 77 Following their entry into force, officially reported emission inventories by 78 parties are increasingly needed to (a) understand and predict source-receptor 79 relationships for such contaminants within both a scientific and regulatory context, and (b) develop sound emission reduction strategies. 80

81 The key objectives of this evaluation have been:

82 (*i*) To identify and discuss specific data needs and requirements regarding
83 emission data for POPs by main users of such information (policy-makers,
84 scientists).

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(*ii*) To compare and contrast policy-driven (official emission data) and
 research-driven emission estimates¹ in terms of major features and selected
 outputs.

(*iii*) To assess temporal trends in relative source contribution for selectedPOPs on the basis official emission data.

90 (*iv*) To discuss if existing emission data include the necessary information
 91 and features for source-receptor relationships to be predicted and understood
 92 and for sound control strategies to be elaborated.

We emphasise that the discussion around official emission data in this study
solely relates to the information submitted by the Parties to the European
Monitoring and Evaluation Programme (EMEP) under the UN/ECE LRTAP
convention by 10th March 2005 (Vestreng et al., 2005).

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98 **2.** Policy and research-oriented features

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100 The features of a specific emission inventory commonly mirrors the specific needs of main user(s) for whom the inventory is targeted. The two key groups 101 102 of users of emission inventories are: (i) policy-makers developing strategies 103 for the reduction of the environmental exposure to pollutants on a regional or global scale, and (ii) scientists studying past, current and future source-104 105 receptor relationships for pollutants on a regional and global scale. It may 106 therefore be useful to distinguish between policy-driven (officially reported 107 emission data by Parties to various international conventions) and researchdriven emission estimates. Policy-driven emission estimates tend to focus on 108 109 identification and control of POP sources, while research-driven emission 110 estimates provide information that helps understanding the impact of POP 111 emissions on the environment and human health. Official policy-driven emission data for POPs are commonly developed and maintained by national 112 emission experts within various environmental protection authorities in the 113 114 Parties to the LRTAP convention (Vestreng et al., 2005). Research-driven 115 estimates are typically developed by groups of international emission experts 116 and target the specific needs of various research projects or environmental 117 assessments. Table 1 contains a simplified classification of various European 118 emission data according to identified policy- and science-oriented features, 119 each which are described and discussed in turn. It should be noted that there 120 are also other emission inventories for POPs available at present. These 121 inventories are not included in Table 1, but they were recently reviewed by 122 Breivik et al. (2004). The main reason for not including all studies conducted 123 so far is the limited geographical scope of some these inventories, i.e. for

¹ Research-driven emission inventories are sometimes referred to as emission expert estimates. It is our opinion that the latter term is inappropriate.

single countries (e.g. Alcock et al. 2001) or for certain parts of Europe only(e.g. Quass et al. 2000; 2004).

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127 **2.1 Policy-oriented features**

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129 An essential policy-oriented feature of an emission inventory is a clear 130 identification of the individual sources and source categories of POP emissions. Official data submitted by the Parties to the LRTAP convention 131 132 (Vestreng et al., 2005) are thought to be superior over the research based emission inventories, because these national emission inventories are 133 134 collected from various industries at national and even international level or 135 they are estimated by emission experts in individual countries. These national 136 experts are expected to know the detailed characteristics in their respective 137 countries concerning relevant activity data, such as usage of individual 138 pesticides and industrial chemicals as well as relevant details concerning the 139 operation of individual plants (e.g. abatement technologies at individual waste 140 incineration facilities).

141 The officially reported POP emissions under the Convention on LRTAP are 142 submitted annually to the UN/ECE Secretariat. It is emphasised that the 143 EMEP emission database is a dynamic database, and the values contained in 144 this database are frequently updated and improved. The requirements for 145 reporting of these data are given in the Emission Reporting Guidelines 146 (UN/ECE, 2002). Emission inventories should be transparent, consistent, comparable, complete, and accurate. The comparability of emission 147 148 inventories is attempted by encouraging all Parties to apply the guidance 149 given in the EMEP/CORINAIR Atmospheric Emission Inventory Guidebook (EEA, 2005). Consistent time series of POPs sectoral and total emissions 150 151 should be reported from 1990 to the latest year (i.e. emission year 2003 for 152 2005 submissions) annually. The emissions should be complete with respect 153 to source categories included. More than 100 source categories are defined in the reporting templates. Gridded data for the priority POPs (PAHs, HCB, 154 PCDD/Fs) are requested every fifth year starting in 1990. For the sake of 155 transparency, all parties are also encouraged to submit an Informative 156 Inventory Report (IIR) together with their emission data in which as a 157 158 minimum, methodologies other than those found in the Guidebook (EEA, 159 2005) and re-calculations should be described and a key source analysis 160 performed. More elaborated IIRs are welcomed, and may include e.g. information on emission factors, emission trend analysis and uncertainty of 161 emission estimates. Since the 2003 trial review of emission data reported to 162 the Convention of LRTAP and under the NEC (National Emission Ceiling) 163 Directive, annual review of emission data has been undertaken and 164 165 documented (e.g. Vestreng et al., 2005). The IIRs have proven to be crucial for a complete understanding of the review results. Like the other 166

167 requirements listed above, the accuracy of the emission data is the responsibility of the parties. The officially reported and review emission data 168 169 are published in the UN/ECE web database. WEBDAB 170 (http://webdab.emep.int/), hosted and maintained by the EMEP Meteorological Synthesizing Centre – West (MSC-W) in Oslo. For most other inventories 171 172 listed in Table 1, the information is usually compiled on an aggregated level 173 taking advantage of aggregated international statistics. It may therefore in 174 many cases be more difficult to take advantage of the "non-official" emission 175 inventories in the policy-oriented context of evaluating possible control 176 measures on a local scale (i.e. for specific emission sources within a country). 177 Prime examples of such aggregated inventories may be the dynamic PCB and 178 PBDE emission mass balance estimates by Breivik et al., (2002a,b) and 179 Prevedouros et al., (2004a), respectively. Other studies seem to rely on 180 official emission data, whenever available, thus only presenting additional 181 emission estimates when there are spatial and/or temporal gaps in the official 182 data (Berdowski et al., 1997; Pacyna et al., 2003; van der Gon et al., 2005).

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185 **2.2 Research-oriented features**

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2.2.1 Spatial coverage

189 For policy-oriented purposes, emission inventories are needed as a useful 190 documentation on the level of reduction of POP emissions agreed within the 191 LRTAP Convention. So far, only a limited number of Parties to the Aarhus 192 POPs protocol within the LRTAP Convention have reported the POP emission data to EMEP. As previously recognised by Berdowski et al. (1997), the 193 194 EMEP emission database has significant gaps in the spatial coverage in 195 reporting of emission data for individual POPs or POP groups (Vestreng et al., 196 2005). As previously mentioned, reliable emission data are essential for 197 efforts aiming to understand and predict source-receptor research 198 relationships POPs (e.g. on a European scale). Therefore, scientists studying 199 these issues request emission data that are complete across respective 200 model domain. In other words, the modeller needs to make sure that there are 201 no regions with no emission data in parts of the spatial domain of the model (if 202 significant emissions are assumed to occur in these regions). Most of the emission data are reported on a country specific level (Table 1), while some 203 emission inventories are further distributed on a grid system, typically a 1° by 204 205 1° grid system for the global emission estimates (Li et al., 2000; 2003, Breivik 206 et al., 2002a) or finer resolution for some of the European emission estimates. 207 For example, the recent European inventory by van der Gon et al. (2005) is presented on a 50 by 50 km² grid system. 208 209

210 2.2.2 Temporal coverage

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212 POP emissions may vary considerably over a variety of time scales (e.g. 213 diurnal, seasonal, inter-annual). In the context of POPs, the key interest is 214 typically to obtain long-term (inter-annual) emission data trends (Table 1), 215 reflecting the potential environmental lifetime of the pollutant of concern. In the 216 case of POPs, this could be decades or even more (e.g. Sinkkonen and 217 Paasivirta, 2000; Wania and Daly, 2002). Many POPs have the potential for 218 undergoing reversible atmospheric deposition (Larsson, 1985; Jeremiason et 219 al., 1994). Atmospheric levels measured today, may thus be a legacy of 220 emissions that had occurred in the past. Several studies have thus attempted 221 to infer the relative importance of primary and secondary emissions in controlling the contemporary atmospheric burden of POPs (e.g. Harrad et al., 222 223 1994; Bailey, 2001; Bidleman and Leone, 2004; Hung et al., 2005; Kurt-224 Karakus et al. 2006).

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226 <u>2.2.3 Speciation</u>

Modellers also need information on the physical and/or chemical speciation of 228 229 individual POP substances (e.g. isomer or congener patterns). This issue may 230 appear to be of particular interest to scientists, but it is argued that it also has 231 important implications for the development of sound control strategies. 232 Information on speciation or congener patterns may be critical because it 233 influences the distance of POP transport within the air masses. PAHs, HCHs, 234 PCBs and PCDD/Fs contain several individual species with different physical-235 chemical properties, environmental degradation rates and thus potential for 236 atmospheric long-range transport (e.g. Beyer et al., 2003). For example Li et al. (2002) have illustrated how β -HCH, unlike α -HCH, exhibited only limited 237 238 propensity for transport into the Arctic in spite of identical chemical formulas 239 and similar emission histories of these two HCHs.

2.2.4 Multimedia features

243 Under certain circumstances and for the more volatile POPs, the multimedia 244 feature of POP releases into the environment could be crucial for 245 understanding atmospheric source-receptor relationships. For example, 246 subsequent transfer of PCBs from contaminated soils and sediments to the atmosphere have been hypothesised to have significant implications for the 247 PCB mass balance (Chiarenzelli et al., 1997). Multimedia emission inventories 248 249 may thus be of particular interest for attempts to evaluate the relative impact 250 of various contaminant pathways to polluted water bodies (e.g. Breivik and 251 Wania, 2002), or studies aiming to evaluate the relative significance of primary 252 anthropogenic (controllable) emissions versus secondary re-emissions from

environmental reservoirs as contaminated in the past (e.g. Harrad et al. 1994;Bailey, 2001; Hung et al. 2005).

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256 2.2.5 Additional data needs

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Some modellers may in addition require more detailed emission input information. These specific needs are typically dependent on the specific geometries of each individual model, and it is therefore considered out of the scope to discuss these specifics in detail. However, it may be mentioned that many of these additional input needs are typically related to physical properties governing emission data (such as emission release heights, particle-size distribution, stack exhaust temperatures and velocities etc).

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266 **2.3 Common data needs**

268 Common to the needs of both policy-makers and scientists is the request for 269 accurate emission information. Information on emission data uncertainty is 270 highly valuable when evaluating the uncertainty of predicted source-receptor relationships as well as the potential risk of choosing erroneous control 271 272 strategies. Little is known about the accuracy of available emission estimates, 273 although a few of the emission inventories listed in Table 1 contain some 274 information on emission data uncertainties. For example, Berdowski et al. 275 (1997) report uncertainty factors for the emission estimates of individual 276 groups of pollutants at European scale (e.g. PAHs: 2-5; PCBs: 2-5 and 277 PCDD/Fs: 5-20), whilst Breivik et al. (2002a,b) and Prevedouros et al. (2004a) 278 present various high and low emission estimates in addition to default 279 estimates. Information on emission data uncertainties in the official emission 280 inventories are contained in the informative inventory reports (IIRs) previously 281 mentioned. IIRs containing information on emission data uncertainties for 282 POPs have recently been submitted by Denmark (Illerup et al. 2005), Finland (SYKE, 2005) and France (CITEPA, 2004). Interestingly, attempts have also 283 284 recently been made to quantify uncertainties in a study of dioxin emission estimates for Central Europe, using Monte-Carlo analysis (Pulles et al. 2006). 285 286

3. A comparison of different inventories for the year 1990

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289 3.1 National totals

One potential approach to evaluate uncertainties of an emission inventory is to compare and contrast emission estimates prepared through different methodologies. Figure 1 compares results from three different emission inventories for selected POPs for the reference year 1990. The reference year 295 1990 is considered to be of specific interest, as the UN/ECE Aarhus protocol 296 on POPs obliges Parties to reduce their emissions of PCDD/Fs, PAHs and 297 HCB below their levels in 1990 (or an alternative year between 1985 and 298 1995). In addition to the EMEP data, Figure 1 also includes results from a 299 comprehensive study by Berdowski et al. (1997), as well as a study by 300 Pacyna et al. (2003). We emphasise that the data from Berdowski et al. 301 (1997) and Pacyna et al. (2003) are based on the guantitative knowledge on POP emissions at the time these studies were published, whereas the EMEP 302 database is frequently updated as new and better information becomes 303 304 available. Thus, differences between these studies may in part be explained 305 by improved characterisation of some specific POP emission sources over the 306 last decade.

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308 <u>3.1.1 PAHs</u>

310 PAHs and dioxins (PCDD/Fs) are the POPs that are most extensively reported to EMEP by the Parties to the LRTAP convention (Figure 1). This 311 312 may not come as a surprise, as the source categorisation and emission inventory methodologies that were originally developed for classical air 313 314 pollutants (EEA, 2005), clearly fit the typical source-profiles of these pollutants better than for pesticides and industrial chemicals. For PAHs, there are both 315 316 official reported EMEP data² and estimates from Berdowski et al. (1997) available for 23 Parties (Fig. 1A). The study by Berdowski et al. (1997) 317 of Borneff six PAHs 318 includes the sum the (benzo[a]pyrene, 319 benzo[*g*,*h*,*i*]perylene, benzo[k]fluoranthene, benzo[b]fluoranthene, fluoranthene and indeno[1,2,3-c,d]perylene), while EMEP request the parties 320 to address four out of these six PAHs (benzo[a]pyrene, benzo[b]fluoranthene, 321 benzo[k]fluoranthene, and indeno[1.2.3-c.d]perylene). A comparison with the 322 323 European emission data from Pacyna et al. (2003) was omitted, as this study 324 considered benzo[a]pyrene only. Differences in speciation are likely to explain 325 in part why the total PAH emissions for the 23 Parties are about three times higher by Berdowski et al. 1997 (in comparison to the EMEP data). However, 326 327 the estimates for individual countries often deviate substantially, and in several cases this difference is more than one order of magnitude (for AT, BG, 328 329 DE, DK, EE, FR, IS, RU)³. Some Parties report higher national emissions than 330 Berdowski et al. (1997) (i.e. BG, CZ, NL). Hence a more limited compound coverage cannot explain the differences between these two inventories. 331

² The EMEP database reports a high value of PAH emissions from "Accidental and Natural Fires" (NFR source category 5E) in GB (about twice as high as the remaining total emissions from the 23 countries included in Figure 1). This value was disregarded in this study, as it is reported as a memo item and should not be included in the national total. The emissions reported from IS are too small (0.087 Mg) to show up in the figure 1A. ³ For interpretation of ISO2 country codes, please see

www.emep.int/grid/country numbers.txt

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333 <u>3.1.2 PCDD/Fs</u>

335 Dioxins are among the POPs that have received a lot of attention, primarily 336 because of their toxicity at very low concentrations. As an unintentional by-337 product of combustion, emissions of dioxins are also expected to take place in 338 all countries⁴. The results of two independent estimates are compared with 339 the EMEP data in Figure 1B. Although there is a fair agreement between 340 these estimates for some countries, large discrepancies (more than 100%) 341 between the EMEP data and one or both independent estimates are evident 342 for BG, CZ, DK, FI, HU, IS, NO, SE, SK.

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344 <u>3.1.3 PCBs</u>

346 Only 9 Parties to the LRTAP Convention have submitted official PCB emission 347 data (greater than zero) for 1990. For the total emissions of PCBs from these 348 9 countries, it can be seen that the estimate by Berdowski et al., (1997) is 349 about twice as high as the EMEP data (Figure 1C). Again, difference in compound coverage within the group of PCBs is an issue that may help to 350 351 explain deviations between these two estimates. The estimates by Berdowski 352 et al. (1997) address total PCBs (i.e. the sum of 209 different compounds) 353 when dealing with emissions from leakage or evaporation - or the sum of six 354 frequently reported congeners (PCB-28, PCB-52, PCB-101, PCB-118, PCB-355 153 and PCB-180) for other emission categories. It is not known which 356 congeners of PCBs are included in the EMEP database. Some countries (e.g. 357 The Netherlands and Spain) are reporting zero PCB emissions to EMEP, and it may be questioned if this is plausible based on the widespread global usage 358 359 of PCBs (Breivik et al., 2002b).

360 The limited availability of emission data for PCBs from various countries 361 indicates that control strategies for PCBs (and other industrial chemicals) may be different in comparison to the PAHs and PCDD/Fs. For example, current 362 363 control strategies in Norway are typically targeted towards the safe collection 364 and destruction of remaining materials and products containing PCBs, rather than based on differences in quantitative source strength with respect to 365 366 atmospheric emissions. Secondly, there are fundamental methodological 367 difficulties addressing emissions of industrial chemicals like PCBs (Breivik et 368 al. 2002b; Prevedouros et al. 2004a). In order to quantify atmospheric 369 emissions of substances that are intentionally produced, consideration must be given to the potential for atmospheric losses that may occur during the 370 371 entire lifecycle of these chemicals (production, transport, storage, use and disposal). Furthermore, there are also fundamental difficulties in determining 372 373 suitable emission factors for PCBs for volatilisation processes (i.e.

⁴ Denmark reports zero emissions of PCDD/Fs in 1990.

evaporation from contaminated products or environmental surface media in 374 375 contact with air). This is because (a) such emissions depend on the 376 equilibrium status between the air and the product / surface media being 377 contaminated with PCBs (e.g. Mackay, 2001), and (b) emissions due to 378 volatilisation are expected to vary significantly in response to temperature 379 changes (e.g. Haque et al. 1974; Haugen et al. 1998; Breivik et al. 2002b). 380 Thus elevated emissions due to volatilisation are expected to take place when 381 "clean" air passes over a contaminated surface at elevated temperatures.

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384 **3.2 Source categories**

386 Overall, the simple comparison presented in Figure 1 provides an idea of the 387 uncertainties in available emission estimates for the individual European 388 countries. A closer look at differences in predicted emissions from various 389 source categories may provide further insight into why there are notable 390 differences between these inventories. Since 2002, emissions reported to the 391 LRTAP Convention, have been reported according to the UN/ECE Emissions reporting Guidelines (UN/ECE, 2002) along the so-called Nomenclature For 392 393 Reporting (NFR) source categories. In order to compare the research-based 394 emission estimates and the EMEP emission data, the NFR categories have 395 been aggregated to the Selected Nomenclature for Air Pollutants (SNAP) 396 according to the corresponding allocation Table III A in the Guidelines 397 (UN/ECE, 2002). For some countries, reported emissions in the SNAP 398 categories have been used directly, since the 1990 emissions were only 399 available in SNAP sectors.

400 Figure 2 shows a comparison of the EMEP data against data presented by 401 Berdowski et al., (1997) and Pacyna et al., (2003). The data are presented at 402 the SNAP Level I for the sake of comparability⁵. Table 2 presents the relative 403 importance of emissions from various SNAP source categories in controlling emissions in selected European countries in 1990. It can be observed that the 404 405 number of countries considered in Figure 2 and Table 2 is lower than the one 406 in Figure 1. This is because some countries only report their national total 407 emissions to EMEP. Also given in the last column in Table 2 are the total 408 emissions in the countries considered.

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410 <u>3.2.1 PAHs</u>

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Table 2 shows that the estimates of PAH emissions by Berdowski et al.,(1997) attribute almost half of the emissions in selected European countries to

⁵ Level 1 is the more coarse aggregation of emission source categories (as opposed to SNAP Levels 2 and 3). A comparison of data at SNAP Level 2 and 3 has not been feasible in this study.

414 SNAP 2 (non-industrial combustion), while the EMEP data attribute about one third of the emissions in this year to SNAP 6 (solvent use). In general, there is 415 416 a very poor correlation between reported emission data from Berdowski et al., 417 (1997) and the EMEP data aggregated to SNAP Level I that are reported for 418 individual countries (Figure 2A). In many cases, the two estimates differ by 419 several orders of magnitude. A closer inspection at the EMEP data in comparison to the data from Berdowski et al. (1997) also indicates that 420 421 important sources in some countries may have been left out in the EMEP 422 data. Prime examples are Estonia and Germany, which both only reports 423 emissions of PAHs from road transport (SNAP 7). It seems safe to claim that 424 for these countries, there is a lack of completeness in terms of source 425 coverage in the EMEP data. As can be seen from Figure 1A, the EMEP data 426 are lower than the data from Berdowski et al., (1997) in 20 out of 23 countries. 427 Figure 2 A also demonstrates this underestimation of EMEP data compared to 428 estimates by Berdowski et al., (1997). Although this may in part be explained 429 by the more limited coverage of PAH species in the EMEP database, it could 430 also be explained by a more complete coverage of emission sources in the 431 study by Berdowski et al., (1997). There are other examples where the data 432 for individual source categories are far higher in the study by Berdowski et al., 433 (1997) in comparison with the EMEP data, but it is not possible to conclude 434 which one is the more correct estimate. A specific example is the emissions of 435 PAHs from non-industrial combustion (SNAP 2) in France. Berdowski et al., (1997) suggest that these emissions are higher than 3000 tonnes, while the 436 437 official EMEP data are almost two orders of magnitude lower (39 tonnes).

439 <u>3.2.2 PCDD/Fs</u>

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A comparison of PCDD/Fs emissions data in EMEP with inventories made by 441 442 Berdowski et al. (1993) and Pacyna et al. (2003) is shown in Table 2 and 443 Figures 2 B,C. Specifically, the total emissions from selected countries are comparable, and all inventories identify waste treatment and disposal (SNAP 444 9) as the primary source category for dioxins on a European scale (Table 2). 445 446 It should be cautioned, however, that waste incineration may not necessarily be a key source of dioxins in all European countries, as previously recognised 447 448 for Belarus (Kakareka, 2002). As can be seen from Table 2, the EMEP 449 database also identifies public power (SNAP 1) and industrial combustion 450 (SNAP 3) as two important source categories for dioxins, while Pacyna et al., (2003) suggest that significant emissions from non-industrial combustion 451 (SNAP 2) as well as production processes (SNAP 4) took place in the 452 453 selected European countries in 1990. These differences may in part be 454 explained by differences in SNAP allocation in individual countries with 455 significant national emissions (Figure 2C). France may serve as an example, although the total emissions in France are suggested to be fairly similar in the 456

reported emission inventories (Figure 1). The EMEP data report the national 457 total emissions to be 1.77 Kg I-TEQ year⁻¹, and the three major source 458 459 categories for which these emissions are attributed are SNAP 1 (47.2%), 460 SNAP 9 (28.0%) and SNAP 3 (20.0%). Likewise, Berdowski et al., (1997) 461 report the national total emissions in France to be 1.64 Kg I-TEQ year⁻¹, for 462 which the key sources are SNAP 3 (54.5%), SNAP 9 (31.4%) and SNAP 2 463 (10.6%). Finally, Pacyna et al., (2003) report the national total emissions in 464 France to be 1.23 Kg I-TEQ year¹, with the key sources being SNAP 9 465 (41.8%), SNAP 2 (30.9%) and SNAP 4 (22.2%).

466 For dioxins, there is an additional concern whether all significant sources have 467 been included in the inventory (completeness). The dioxin emissions from the 468 open burning of household waste have received considerable attention in 469 recent years (e.g. Lemieux et al., 2000/2004; Wevers et al., 2004). However, 470 reliable estimates of the relative importance of such emissions are considered 471 difficult because of the lack of reliable activity data and emission factors 472 related to open burning. 473

- 474 <u>3.2.3 PCBs</u>
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476 Only six European countries have reported official emission data for PCBs by 477 source category for the year 1990. The total PCB emissions from these six 478 countries amount to 10.4 tonnes according to EMEP, and 27.1 tonnes 479 according to Berdowski et al., (1997). In the case of the EMEP data, 78.6% of 480 these emissions are attributed to SNAP 6. According to the explanatory notes 481 from United Kingdom (which reported the highest PCB emissions among the 482 countries listed in Table 2 – see also Figure 1), this lumped source category 483 includes emissions from capacitors and transformers and the use of halogenated chemicals in the chemical industry. Berdowski et al., (1997) also 484 485 report that most of the PCB emissions were attributed to electrical equipment, 486 but did not assign these emissions to any existing SNAP source category 487 (Table 2).

489 4 Temporal changes in key source categories

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490 491 It is argued that official emission data should be the preferred choice of 492 emission information, as official estimates seem to be most suitable for 493 evaluation of emission reductions. Table 3 presents a key source analysis for 494 selected POPs according to the EMEP data alone. The determination of key

495 sources is a fairly simple ranking technique, which is considered useful to identify the importance of various sources in drawing conclusions about the 496 497 emission current levels and emission trends (e.g. Rypdal and Flugsrud, 2001; 498 Rypdal, 2002). Inventory improvements may then be directed towards the key 499 sources that have been identified, although there is a general concern if all 500 key sources are included in official national emission inventories (see 501 discussion above). For simplicity, only the top ten source categories are listed 502 in Table 3 for those pollutants that have more than ten source categories 503 contributing to 95% of the total emissions. It should be observed that the 504 number of countries included in the comparison is rather limited. This is 505 because we only included Parties for which data in NFR (Nomenclature for Reporting) format are available (Vestreng et al., 2005) for the years 506 507 considered in this study. Table 3 includes data for both 1990 and 2003 in 508 order to evaluate potential temporal changes of the key sources. The reported 509 emission reductions in selected European countries from 1990 to 2003 are 510 ~80% in the case of PCDD/Fs and PCBs, whereas a reduction of ~47% is suggested for PAHs in the same period (2nd last row of Table 3). Only a few 511 512 Parties report increasing emission from 1990 to 2003 (BE, DK and IS in the 513 case of PAHs, and FR in the case of PCBs).

The frequent assignment of POP emissions to "other" source categories as well as the limited number of sources listed, serve to illustrate that the official reporting scheme (originally developed for classical air pollutants) may be considered less suitable when applied to industrial chemicals and pesticides. An extension of the NFRs to increase the completeness and transparency in the reporting of POPs should therefore be considered when the UN/ECE Emission Reporting Guidelines are to be revised in 2007.

521 For PAHs, various (other) processes in the chemical industries and metal 522 production were the two key sources in 1990 (Table 3). By 2003, residential 523 plants are suggested to be the key source of PAHs, contributing almost a third 524 of the total emissions from the European countries included in Table 3. The 525 data indicate that a significant shift has occurred over the time, in which 526 emissions from residential plants have increased in relative importance on the 527 expense of various industrial emissions. Likewise, residential plants are also 528 suggested to be the most significant source of dioxin emissions in 2003, 529 following stringent emission controls in other sectors (e.g. waste incineration 530 from 1990). For PCBs, only five source categories are addressed and only 531 two countries are included (GB, FR)⁶. The total PCB emissions from these two countries are dominated by emissions from GB (99% and 94% in 1990 532 533 and 2003, respectively), and other processes in the chemical industries (i.e. 534 emissions from capacitors and transformers in the case of G.B.) are attributed 535 as the key source in both years.

537 5 Final remarks

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⁶ SE reports minor national emissions of PCBs, but for "national navigation" only. In addition, SE also reports PCB emissions from "international navigation".

539 While a number of efforts have been made to improve emission inventories 540 needed for better understanding major environmental problems, such as 541 stratospheric ozone depletion, climate change and acid deposition, inventories 542 for POPs have for many years been considered unreliable and inaccurate 543 (Graedel et al., 1993; Pacyna and Graedel, 1995). A number of studies have 544 concluded that emission data are frequently the most uncertain input 545 information that determines the overall uncertainty of model predictions for 546 POPs (Vallack et al., 1998; Cohen et al., 2002; Malanichev et al., 2004). In 547 fact, it has often been claimed that the emissions of POPs remain the least 548 understood part of the research on overall distribution and fate of these 549 chemicals in the environment (Vallack et al., 1998; Wania and Mackay, 1996; 550 Jones and de Voogt, 1999; Breivik and Alcock, 2002). The simple comparison 551 of emission data in this study illustrates some of the difficulties with current 552 emission data for POPs, and in comparing and contrasting emission data of 553 different origin.

554 At present, it is not possible to firmly conclude which of the available emission 555 inventories are the most accurate. Even so, it is suggested that individual 556 countries may take further advantage of the research-based emission 557 inventories (i.e. non-official data) and use this information in search for 558 improvement of their official national emission inventories. For example, 559 research-based emission inventories may assist in identifying and filling gaps 560 with respect to some significant source categories that may have been 561 ignored or missing in the official emission data. The lack of consideration of 562 other PAH sources than road transport in the German and Estonian emission 563 inventories may serve to illustrate this point. It is also concluded that the non-564 official data are at elevated risk of providing erroneous estimates, due to a 565 lack of knowledge on specific factors or control strategies that may have been 566 implemented in various individual countries.

567 Whilst official emission data should be the preferred choice of emission 568 information, it is shown that the EMEP data contain incomplete information on spatial, temporal, and speciation features. This generally mitigates the use of 569 570 official emission data in the context of source-receptor modelling on a regional 571 (e.g. European) scale. For this reason, modellers may still have to rely on 572 research-driven estimates, which have typically aimed to present the more 573 general picture of emissions of individual POP substances in quantitative 574 terms (e.g. Breivik et al., 2002a,b; Prevedouros et al., 2004a).

575 The upcoming revision of the UN/ECE Emissions Reporting Guidelines and 576 future studies on emission inventories of POPs shall recognise and address 577 more clearly the specific features of POP emission inventories related to their 578 accuracy and completeness⁷. This consideration is needed for both scientific 579 and regulatory purposes of developing emission inventories of POPs.

⁷ The interested reader may note that a new report on the EMEP emission data has been issued since the submission of this manuscript (Vestreng et al. 2006)

580 Further development of official emission data is also dependent on the 581 outcome of future research on the generation of POPs during various 582 anthropogenic and natural processes. This research should focus on the 583 improvement of methodological approaches leading to the establishment of 584 more accurate methods of POP emission estimates.

585 The outcome of this study also calls for further efforts on the standardisation 586 of POP emission inventories and data parameters. This is an important issue 587 that still needs further consideration for future comparative studies and interregional decision-making on a regional European basis. This study may in this 588 589 respect also have potential implications for relevant regional activities taking 590 place on other continents and even globally (i.e. work under the Stockholm 591 Convention on POPs; UNEP, 2001). As for this latter agreement, an important 592 step for dioxins and furans has already been made. For these chemicals a 593 standardised toolkit for identification and quantification of dioxin and furan 594 releases have been issued to assist countries to establish emission 595 inventories (UNEP, 2005). Use of this guidebook among countries is thus 596 expected to contribute to consistent emission data over time and between 597 countries.

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603 **References**

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854 **Biographical note**

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897	Figure captions
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- Figure 1 Comparison of different emission estimates for selected POPs in
 1990. The data are sorted from EMEP high to EMEP low. The
 numerical values are reported in Vestreng et al., (2005)⁸.
- Figure 2 Comparison of emission estimates from Berdowski et al., (1997)
 and Pacyna et al., (2003) versus EMEP data at SNAP Level 1 for
 the reference year 1990. A) PAHs (unit in tonnes year⁻¹), B) and C)
 PCDD/Fs (unit in g I-TEQ year⁻¹).

⁸ The data reported by Berdowski et al. (1997) sometimes refer to officially reported data submitted for a different reference year than 1990. Data for PAHs and PCBs from Germany refer to data for 1985-1990, data from Luxembourg and the Netherlands refer to data from 1993, while data from Sweden refer to the years 1987 and 1991.

Table 1. Simplified classification of selected emission inventories for POPs
 with European coverage, according to policy- and science-oriented
 features (*see text for discussion*). A parenthesis indicates that some
 information is available on the subject. NA = Not applicable.

	Policy- oriented features	Resea	Common			
	Source identification	Spatial (by country)	Temporal (inter- annual)	Speciation	Multimedia	Uncertainty
EMEP (official data)	+	(+)	(+)	(+)	-	(+)
Berdowski et al., 1997	(+)	+	-	-	-	(+)
Pacyna et al., 2003	(+)	+	+	(+)	-	-
Prevedouros et al., 2004a	(+)	+	+	+	-	(+)
Van der Gon et al., 2005	(+)	+	+	(+)	-	-
Li et al., 2000/2003	(+)	+	+	+	-	-
Breivik et al., 2002a,b	(+)	+	+	+	-	(+)
Bailey (2001)	(+)	-	-	NA	-	(+)

Table 2. Relative importance of emissions from various SNAP source categories^{A)} in controlling emissions in selected European 945 countries^{B)} in the reference year 1990 (%). The latter column gives the total emissions in the European countries considered (Units are thousand tonnes per year for PAHs, kg I-TEQ per year for PCDD/Fs and tonnes per year for 947 PCBs), NI = No information. 948

			SNAP (% of Total)							Total				
		1	2	3	4	5	6	7	8	9	10	11	Other	
Hs	EMEP (2005)	2.7	20.5	6.4	19.9	<0.1	34.4	10.6	1.5	2.1	2.0	<0.1	NI	3.5
٩d	Berdowski et al. (1997)	0.3	45.2	1.3	11.5	0	28.8	11.2	1.1	<0.1	0	0.4	NI	12.5
DD/Fs	EMEP (2005)	20.1	18.2	19.9	12.2	0.1	0.1	1.3	0.9	25.6	0.8	0.8	NI	8.4
	Berdowski et al. (1997)	7.1	19.7	15.7	23.0	<0.1	<0.1	1.0	0.5	33.0	0	<0.1	NI	7.5
РС	Pacyna et al. (1997)	5.7	30.0	6.0	26.4	0	<0.1	1.9	0.2	29.5	0	0.3	NI	8.9
PCBs	EMEP (2005)	6.5	2.6	2.4	5.7	0	78.6	1.4	0.2	2.6	0	0	NI	10.4
	Berdowski et al. (1997)	3.1	0.5	0.4	1.3	0	0	0.3	0	0.1	0	0	94.2	27.1

Definition of SNAP source categories: 1. Combustion in energy and transformation industries; 2. Non-industrial combustion; A) 3. Combustion in manufacturing industry; 4. Production processes; 5. Extraction and distribution of fossil fuels and geothermal energy; 6. Solvent and other product use; 7. Road transport; 8. Other mobile sources and machinery; 9. Waste treatment and disposal; 10. Agriculture; 11. Other sources and sinks. Other: Electrical equipment.

PAHs: NL, BG, GB, BE, ES, PL, IT, FR, SK, SE, SI, RU, AT, FI, HR, NO, DK, MD, DE, EE, IS. PCDD/Fs: FR, GB, DE, RU, B) BÉ, BG, PL, IT, CH, SK, ES, AT, NO, SE, LU, FI, IS. PCBs: GB, PL, SI, BG, SK, FR.

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Table 3. Key Source Analysis for selected POPs in 1990 and 2003 in European countries for which official (EMEP) data for both 959 years are available^{A)}. The numbers in parenthesis give the relative contribution to total emissions, whilst the total emissions are presented in the 2nd last row. The last row gives the number of source categories not included, as only the 961 top ten source categories are listed. 962

PA	Hs	PCD	D/Fs	PCBs		
1990	2003	1990	2003	1990	2003	
Other industrial	Residential plants	Waste incineration	Residential plants	Other industrial	Other industrial	
Metal production	Solvent & other product	Public electricity & heat	Waste incineration	Metal production	Waste incineration	
(21.2)	use^{B} (17.9)	prod. (25.1)	(17.9)	(6.9)	(10.9)	
Solvent & other product use ^{B)} (12.7)	Other industrial processes (12.3)	Other, waste (9.9)	Public electricity & heat prod. (14.7)	Waste incineration (2.6)	Metal production (9.5)	
Residential plants (8.0)	Metal production (9.3)	Metal production (9.3)	Iron and steel (10.4)		Public electricity & heat prod. (8.1)	
Road transport	Road transport,	Iron and steel	Metal production		Iron and steel	
passenger cars (5.8)	passenger cars (4.0)	(8.6)	(9.8)		(1.7)	
Field burning of agric. waste (4.2)	Non-ferrous metals (3.1)	Residential plants (3.2)	Other, manufacturing industries & construction (7 0)			
Road transport, heavy duty vehicles (3.2)	Road transport, heavy duty vehicles (2.8)	Non-ferrous metals (2.3)	Non-ferrous metals (5.4)			
Paint application (1.7)	Field burning of agric. waste (2.2)	Commercial / insti- tutional plants (2.1)	Solvent & other product use ^{B)} (2.1)			
Non-ferrous metals (1.5)	Waste incineration (1.9)	Other, manufacturing industries & construction (1.6)	Other, waste (1.8)			
Solid fuel transformation (1.5)	Road transport, light duty vehicles (1.7)	Field burning of agric. waste (1.4)	Petroleum refining (1.5)			
2.4 ktonnes year-1	1.3 ktonnes year ⁻¹	4.9 Kg I-TEQ year ⁻¹	1.0 Kg I-TEQ year-1	7.2 tonnes year-1	1.5 tonnes year ⁻¹	
41	41	2	5	0	0	

A) The following countries are included in the key source analysis: PAHs: AT, BE, DK, FR, IS, NL, NO, ES, SE, GB. PCDD/Fs: AT, BE, FR, IS, NL, NO, ES, SE, GB. PCBs: FR, GB. B) Including products containing POPs.

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Figure 1



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970 Figure 2



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