

Air concentrations and deposition of chlorinated dioxins and furans (PCDD/F) at three high Alpine monitoring stations: Trends and dependence on air masses

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Abstract

Ambient air concentrations and bulk deposition of polychlorinated dibenzo-*p*-dioxins and dibenzofurans (PCDD/F) were monitored at three high Alpine stations in Germany (Schneefernerhaus/Zugspitze: 2650 m a.s.l.), Switzerland (Weißfluhjoch: 2663 m a.s.l.), and Austria (Sonnblick Observatory: 3106 m a.s.l.) from 2005-2018. High-volume active air samplers were selectively in operation on a daily basis according to predicted source-regions of air masses. Air concentrations showed only small differences between the three sites and were one or two orders of magnitude lower than in urbanized areas in central Europe. Calculation of source contribution revealed waste incineration, heating, and coal combustion as major sources of PCDD/F in ambient air. Air masses from the northeast were characterized by higher concentrations of PCDD/F than those from northwest and south. However, air masses from the northeast source region are significantly less frequent at the three Alpine stations than air masses from the other source regions. Only at Schneefernerhaus, the majority of congeners exhibit a clear decreasing temporal trend in ambient air concentrations. Deposition rates of PCDD/F measured at the three stations were almost as high as at monitoring stations in urban agglomerations. Despite lower precipitation rates in the central Alps, deposition of PCDD/F was higher at the Sonnblick Observatory than in the Northern Alps (Schneefernerhaus). Bulk deposition for Weißfluhjoch and Schneefernerhaus exhibit a decreasing temporal trend.

Keywords: Active air sampling, Alps, Bulk deposition, EPA-PMF, FLEXPART, long range atmospheric transport (LRAT), MONARPOP, PureAlps

1. Introduction

Meteorological conditions which are characteristic for mountain ranges, such as low temperatures, high levels of precipitation and pronounced valley winds, cause high reception of persistent organic pollutants (POPs) (Daly and Wania, 2005; Wang et al., 2007; Offenthaler et al., 2009). This is particularly the case

57 for the Alps, which are surrounded by regions with pronounced industrial activities and a high
58 population density (Iozza, 2010). According to van der Gon et al. (2007), total emissions of
59 polychlorinated dibenzo-*p*-dioxins and dibenzofurans (PCDD/F) in 2000 amounted to 11.7 kg toxicity
60 equivalents (TEQ) yr⁻¹ for Europe (UNECE official data and estimates). The emissions were 0.050,
61 0.017, 0.744, 0.406, 0.560, 0.245 and 0.027 kg TEQ yr⁻¹ for Austria, Switzerland, Czech Republic,
62 Germany, France, Italy, and Slovenia, respectively.

63
64 Within the international monitoring project MONARPOP (“Monitoring Network in the Alpine Region
65 for Persistent and other Organic Pollutants”), beginning in 2005, unprecedented measurements of
66 chemicals which belong to the “Dirty Dozen” were performed in mountainous ecosystems in several
67 regions of Austria, Italy, Slovenia, Switzerland, and Germany. In consequence, numerous scientific
68 publications drew the attention to the Alpine region as a sink for POPs (Finizio et al., 2006; Belis et al.,
69 2009; Jakobi et al., 2015).

70
71 Altitudinal profiles of POPs in soils and plant tissue of alpine ecosystems showed that for some
72 compounds (e.g. organochloropesticides) there is an increase from valley floor to high elevations due to
73 decreasing air temperatures (Kirchner et al., 2009). On the basis of horizontal transect measurements
74 through the Alps, Offenthaler et al. (2008) found out that concentration levels of PCDD/F,
75 polychlorinated biphenyls (PCB) and polycyclic aromatic hydrocarbons (PAH) in humus and needles
76 are higher in the northern and southern fringes of the Alps than in central Alps.

77
78 Continuous measurements of air concentrations and bulk deposition at high elevation sites at Hoher
79 Sonnblick, Weißfluhjoch and Zugspitze (Schneefernerhaus) were an additional part of the MONARPOP
80 project which are continued by the projects PureAlps in Austria and Bavaria/Germany. Hence, the time
81 series presently ranges from 2005 to 2018 (only at Weißfluhjoch measurements ended in 2013).
82 Evaluation of this comprehensive time-series on POPs started with a focus on organochloropesticides
83 (OCPs). The respective results of deposition rates and air concentrations of OCPs have been published
84 recently (Jakobi et al., 2015; Kirchner et al., 2016).

85
86 The present paper extends the evaluation of the MONARPOP/PureAlps data and the determination of
87 PCDD/F in air and bulk deposition at the three high elevation sites. Deposition of PCDD/F can occur in
88 dry gaseous, dry particulate and wet forms (Lohmann and Jones, 1998). Several studies at sites ranging
89 from mountains (Suryani et al., 2015), polar regions (Oehme et al., 1996; Kallenborn et al., 2012;
90 Kallenborn et al., 2013), sparsely populated rural areas (Hovmand et al., 2007) to more polluted urban
91 areas (Wallenhorst, 1997; Harrad, 2010) have been performed and give a range of atmospheric
92 concentrations and deposition rates of PCDD/F for various parts of our globe. The present study
93 supplements these studies with data from the Alpine region.

94
95

96 **2. Materials and methods**

97
98 2.1. Sampling sites and meteorological conditions

99
100 Sampling has been performed at three mountain summit stations: In Bavaria/Germany at
101 Schneefernerhaus (UFS: 2650 m a.s.l.; 47.42°N; 10.98°E) in close vicinity of the highest mountain of
102 Germany (Zugspitze), in Austria at Sonnblick Observatory (SBO: 3106 m a.s.l.; 47.05°N; 12.96°E), and
103 in Switzerland at Weissfluhjoch (WEI: 2663 m a.s.l.; 46.83°N; 9.81°E). The horizontal distances
104 between WEI, UFS, and SBO to the nearest settlement areas as possible sources of PCDD/F are 4, 10,
105 and 20 km, respectively. The annual precipitation amounts monitored at UFS, SBO and WEI were 2551,
106 1829 and 1379 mm during 2005-2013; the amounts of precipitation show a certain dependency from the
107 distance from the northern edge of the Alps (UFS: 20 km; WEI: 60 km; SBO: 70 km) and from the
108 altitude. Jakobi et al. (2015) have provided more detailed information about the sites and the
109 meteorological conditions during 2005-2013.

110
111

112 2.2. Sample collection

113
114 Air sampling of polychlorinated dibenzo-*p*-dioxins and dibenzofurans (PCDD/F) at the three stations
115 was performed by using multichannel high-volume samplers (HVS; Dioxin Monitoring Systems:
116 Kottlingbrunn, Austria; Kroneis: Vienna Austria and Digitel: Hegnau, Switzerland) with a flow rate of
117 120 L min⁻¹ based on the German guideline VDI 3498-1. The HVS were equipped with a switching unit
118 of heated magnetic valves to select one of the 4 filter cartridges, an Ethernet interface and an extended
119 control unit to forward and process control and status-information (Offenthaler et al., 2009). The filter
120 cartridges were equipped with heated inlet tubes to avoid freezing of the glass fiber filters of the filter
121 cartridges. ¹³C₁₂-labeled 1,2,3,4-TCDD was added as sampling standard prior to exposure of the
122 cartridges. The HVS had to operate for 3 months to collect detectable amounts within the cartridges.

123
124 Until 2015, air sampling was designed to distinguish three predefined source regions: northwest (NW),
125 northeast/east (NE) and southwest/south/southeast (S) (Fig. 1). A fourth region of nominally undefined
126 origin (UD) was assigned for air masses which remained less than three days over any of the three source
127 regions (Kirchner et al., 2016). Such air masses mostly came directly from the Atlantic Sea. The
128 selection of the four sampling filters was governed by daily trajectory forecasts calculated using the
129 FLEXPART model developed by NILU (Stohl et al., 2005); the samplers were programmed on a daily
130 basis by the Austrian Central Institute for Meteorology and Geodynamics (ZAMG), Regional Office
131 Salzburg. After 2015, air sampling has been performed without differentiating the four source regions.

132
133



135
136 Figure 1: Directional selectivity of active air sampling (Offenthaler et al., 2009)

137
138
139 To sample atmospheric bulk deposition under the extreme climatic conditions at the Alpine sites, new
140 funnel-adsorber samplers have been developed. Bulk deposition samplers according to German
141 guideline DIN 19739-1 were equipped with additional isolation, heated glass funnels to melt snow, and
142 heated cartridge chambers to avoid frost shattering. Precipitation is collected in a funnel with a sampling
143 area of 0.053 m², melts and subsequently runs through a glass cartridge, which is filled with styrene-
144 divinylbenzene copolymer resin as a hydrophobic adsorber (XAD-2 resin, Supelpak-2SV, Supelco).
145 Detailed description can be found in Offenthaler et al. (2009) and Jakobi et al. (2015). PCDD/F
146 deposition rates are determined by both the amount of precipitation and its PCDD/F content.

147
148
149 2.3. Substances, chemical analysis and quality assurance/quality control

150
151 We monitored 7 toxic 2,3,7,8-chlorosubstituted congeners of the PCDD and 10 toxic 2,3,7,8-
152 chlorosubstituted congeners of the PCDF, the sum of tetra- (TCDD and TCDF), penta- (PeCDD and
153 PeCDF), hexa- (HxCDD and HxCDF) and heptachlorinated (HpCDD and HpCDF) and octachlorinated

154 (OCDD and OCDF) homologues. The total sum of PCDD/F and the toxicity equivalent for humans
155 (TEQ) based on the toxic equivalency factors (TEF) according to NATO/CCMS 1988 (I-TEF) and the
156 World Health Organisation (WHO) 2005 (Van den Berg et al. 2006) were calculated on a lower bound
157 basis from thereof.

158
159 All sample materials (XAD-2 resins as adsorber material from deposition samplers, glass fiber filters
160 and PUF plugs from active air samplers) were spiked with 17 ¹³C₁₂-labeled 2,3,7,8-substituted PCDD/F
161 congeners prior to extraction (Cambridge Isotope Laboratories Inc., Tewksbury, Massachusetts, USA).
162 ¹³C₁₂-labeled 1,2,3,4-TCDD was used as sampling standard to have all 17 ¹³C₁₂-labeled 2,3,7,8-
163 substituted PCDD/F congeners available for recovery correction. For the active air samples extraction
164 was carried out in a Soxhlet extractor for 12 hours using toluene. For deposition samples, the outside of
165 each cartridge was cleaned using toluene wetted pads. The so cleaned cartridges containing the sample
166 material were extracted as a whole in a Soxhlet extractor equipped with a Dean-Stark water separator
167 for three days.

168
169 The clean-up of the extracts was done by a semiautomatic three step liquid column procedure
170 [H₂SO₄/Celite, mixed layer column (acidic silica, neutral silica, basic silica) and alumina B column],
171 also including the separation of PCDD/F from PCB. The extracts were reduced to 50 µl by a gentle
172 stream of nitrogen, ¹³C₁₂-labeled 1,2,3,4-TCDF was added as recovery standard prior to GC/HRMS
173 measurement. The GC/HRMS measurement was carried out on a MAT 95XP high resolution mass
174 spectrometer (Thermo Finnigan, Bremen, Germany) connected with two Trace GC Ultra gas
175 chromatographs (Thermo Scientific). One GC was equipped with a DB-DIOXIN capillary column
176 (Agilent J&W; length: 60 m, Ø: 0.25 mm, film thickness: 0.15 µm), the other one with a DB-5 capillary
177 column (Agilent J&W; length: 60 m, Ø: 0.25 mm, film thickness: 0.25 µm), respectively. Each sample
178 was analyzed with both GC columns to achieve a good separation of all 2,3,7,8-substituted congeners
179 and the possibility to calculate the sum of homologue groups. The measurements were carried out at a
180 mass resolution of 8,000 to 10,000 in the single ion monitoring (SIM) mode with the two most intensive
181 masses of the molecule ion cluster of each homologue group of the native and ¹³C₁₂-labeled PCDD/F.
182 Quantification was done using isotope dilution method with the corresponding ¹³C₁₂-labeled extraction
183 standard for each 2,3,7,8-substituted congener according to quality criteria given in ISO 16000-14. For
184 the sum of the homologue groups the area of every peak within the retention-time window which fulfils
185 the quality criteria was summed up and quantified using isotope dilution method with one corresponding
186 ¹³C₁₂-labeled extraction standard for each homologue group.

187
188 Field blank samples have been analyzed for ambient air as well as for deposition measurements. For
189 ambient air measurements filter cartridges have been exposed in a meteorological cabin made of
190 untreated wood to ambient air without suction for the same period as the cartridges at the active sampler.
191 For deposition measurements two types of field blanks have been used during the project. In the early
192 stage the glass cartridges filled with XAD-2 resin were exposed open to ambient air but protected from
193 precipitation in a meteorological cabin, to estimate the passive sampling capacity. In that type of field
194 blanks no detectable concentrations could be found for PCDD/F. Therefore, the type of field blanks has
195 been changed to glass cartridges filled with XAD-2 resin, which were sent to the sampling site, stored
196 closed in the heated sample chamber of the deposition sampler together with the sampled cartridge. At
197 the end of the sampling period, the field blank cartridges were sent together with the deposition samples
198 back to the laboratory. Blank values were always below 10% of the lowest measured concentrations of
199 each batch and could therefore be accepted without subtraction according to ISO 16000-14.

200
201 The accuracy of the analytical method has been determined with regularly participation in proficiency
202 tests (PT). The standard deviation of the method was determined by parallel sampling of ambient air at
203 the same sampling site and separate determination. For results calculated as TEQ the standard deviation
204 amounts to < 10%. For individual congeners the mean standard deviation is in the range of 25%, near
205 LOQ the standard deviation can raise up to 60% (Moche and Thanner, 1996).

206 Due to the used sampling procedure as described above the sampled air volumes are strongly depending
207 on the weather conditions and thus differ from a few hundred cubic meters to several thousand cubic
208 meters for the individual samples. For this reason the limits of quantification (LOQs) of the method also

209 vary in a wide range from 0.001 to 0.08 fg m⁻³ for active air sampling. For deposition samples LOQs are
210 in the range from 0.001 to 0.07 pgm⁻²d⁻¹ for the individual congeners.

211
212 Breakthrough experiments have been carried out in the way that the two PU-foams were analysed
213 separately for some cartridges. PCDD/F could not be found in significant concentrations on the second
214 PU-foam.

215
216 Standard statistical methods were applied for the presentation of data (range, median). Pearson's
217 correlation and a t-test were used for trend analysis of compounds; p=0.05 and p=0.001 denoted
218 statistical significance (Anderson and Finn, 1996).

219
220

221 3. Results and discussion

222
223 3.1. Data availability and frequency of air masses

224
225 In our study, 19 (WEI) to 31 (UFS, SBO) sampling periods have been evaluated. In the case of active
226 air concentration monitoring, mean data availability (in percent of length of the entire time span) was
227 82%, 86%, and 90% for UFS, SBO, and WEI, respectively. Periods in which data availability due to
228 station or device problems was lower than 40% were excluded (Kirchner et al., 2015). Deposition data
229 were available in 97% (UFS), 94% (SBO), and 81% (WEI) of the entire time span.

230
231 Weather conditions at the three sites were mainly characterized by air masses originating in the
232 Mediterranean region (S) (32-35%) and the NW region (27-32%). Relatively few air masses originated
233 from the NE source region (11-15%), whereas the frequencies of air masses with the unspecific origin
234 UD were 22-25%. The accuracy of forecasts, which had been evaluated for the first three years of
235 measurements by *post hoc* inspection of the trajectories of air actually collected by each filter unit, was
236 high for NE and NW, but lower in the case of prefrontal southwesterly winds and foehn events (Kaiser,
237 2009). Additional information about the trend of air masses arriving at the three sites can be found in
238 Kirchner et al. (2015).

239
240

241 3.2. Mean air concentrations and seasonality

242
243 The concentrations of 17 2,3,7,8-substituted congeners, the sums of TCDD/F, PeCDD/F, HxCDD/F and
244 HpCDD/F and the total sum of PCDD/F are shown in Table 1. Additionally, the toxicity equivalents
245 (TEQ) according to NATO/CCMS (1988, I-TEF) and the World Health Organisation (WHO-2005) are
246 reported (Van den Berg et al., 2006). The concentrations were continuously normalized to 0 °C and
247 1013 hPa by the HVS according to the Avogadro equation (Ackermann and Knox, 2006). The overall
248 concentrations of singular periods were calculated by weighting concentrations by the relative
249 abundance of air masses from each source region. Based on calendar date, each period was attributed to
250 the colder and warmer half of the year and calculated winter-to-summer ratios.

251
252 Regarding the entire period between 2005 and 2013/2018, dioxin concentrations in air were found to be
253 6.0-195 fg m⁻³ (sum PCDD/F) and 0.001-6.0 fg I-TEQ m⁻³. For all three sites, PCDD ranged between 2
254 and 164 fg m⁻³, and PCDF between 0.3 and 171 fg m⁻³. The concentration ranges and medians of all
255 congeners, the corresponding sums and TEQ values were similar for the three sites and did not differ
256 significantly between them. However, some congeners were frequently below the limit of detection.
257 After omitting the outliers, only OCDD at SBO and UFS was significantly higher than at WEI (p<0.05).

258
259 At our Alpine sites, the PCDD/F concentrations were up to 1-2 orders of magnitude lower than those
260 monitored in urban areas or emission centers. In the Rhine-Ruhr conurbation, annual means of PCDD/F
261 concentrations were in the range of 15-50 fg I-TEQ m⁻³ in the period 2000-2010 (Bruckmann et al.,
262 2013). During a monitoring program (1997-1999) performed in different urban regions of Austria
263 (Moche and Thanner, 2003), annual means varied from 36 fg I-TEQ m⁻³ (Vienna) to 237 fg I-TEQ m⁻³
264 (Donawitz). In urban areas of Baden-Württemberg (Germany), average dioxin concentrations ranged

265 from 53-99 fg I-TEQ m⁻³, whereas the average PCDD/F concentrations in rural areas were found to be
 266 14-27 fg I-TEQ m⁻³ in mid-nineties (Wallenhorst et al., 1997). Around a steel plant area in northeast
 267 China, the air concentrations and WHO-TEQ values of PCDD/F were in the range of 94-4944 fg m⁻³ and
 268 3-247 fg m⁻³, respectively (Li et al., 2010). Within the National Dioxin Air Monitoring Network
 269 (NDAMN), PCDD/F concentrations in air were in the range of 6-15 fg TEQ m⁻³ in rural areas, and of
 270 0.1-3 fg TEQ m⁻³ in remote areas of the USA (EPA, 2013). Rural background locations in
 271 Catalonia/Spain exhibit geometric mean concentrations of 16 fg TEQ m⁻³ (Parera et al., 2018). At a
 272 remote site (Zöbelboden) of the Austrian National Park “Kalkalpen” at 900 m a.s.l., the three-year mean
 273 from 1997 to 1999 was 2.7 fg I-TEQ m⁻³ for summertime and 4.4 fg I-TEQ m⁻³ for wintertime (Moche
 274 and Thanner, 2003). At an urban background and one rural site in Bavaria medians of PCDD/F
 275 concentrations were 6.7 and 9.0 fg WHO-TEQ m⁻³ in summertime and 33 and 35 fg WHO-TEQ m⁻³ in
 276 wintertime in 2002-2004 (Bayerisches Landesamt für Umwelt, 2006).
 277 Results from Ny-Alesund, Spitsbergen (Norway), show that OCDD and OCDF concentrations in the air
 278 during summer 1995 were 4.4 and 3.8 fg m⁻³, respectively and the sum of PCDD/F concentrations ranged
 279 between 67 and 105 fg m⁻³ in spring and summer samples (Schlabach et al., 1996).

282 Table 1: Concentrations of PCDD/F measured with HVS at Hoher Sonnblick (SBO) in 2005-2018,
 283 Weissfluhjoch (WEI) in 2005-2013 and Schneefernerhaus/Zugspitze (UFS) in 2005-2018: Ranges
 284 (minimum-maximum), medians and winter to summer ratios calculated on the basis of medians
 285 (including outliers)

Compound	SBO			WEI			UFS		
	Range fg m ⁻³	Median fg m ⁻³	Winter to summer ratio (medians) *p<0.05	Range fg m ⁻³	Median fg m ⁻³	Winter to summer ratio (medians) *p<0.05	Range fg m ⁻³	Median fg m ⁻³	Winter to summer ratio (medians) *p<0.05
2,3,7,8-TCDD	n.d.-n.d.	n.d.	-	n.d.-0.4	n.d.	-	n.d.-0.1	n.d.	-
1,2,3,7,8-PeCDD	n.d.-0.3	n.d.	2.5	n.d.-0.2	n.d.	-	n.d.-0.2	n.d.	-
1,2,3,4,7,8-HxCDD	n.d.-3.2	n.d.	2.5	n.d.-0.8	n.d.	-	n.d.-0.5	n.d.	1.6
1,2,3,6,7,8-HxCDD	n.d.-3.2	n.d.	0.9	n.d.-1.0	0.1	0.6	n.d.-1.4	n.d.	1.2
1,2,3,7,8,9-HxCDD	n.d.-1.8	n.d.	1.0	n.d.-0.9	n.d.	-	n.d.-0.7	n.d.	0.8
1,2,3,4,6,7,8-HpCDD	n.d.-16.0	2.9	0.8	n.d.-9.7	1.9	1.6	n.d.-13.4	1.9	1.3
OCDD	4.1-56.7	9.3	1.0	n.d.-148	7.5	1.1	n.d.-40.0	8.7	1.5
Sum TCDD	n.d.-11.8	2.0	1.3	n.d.-8.4	1.4	1.2	n.d.-21.2	1.5	2.1*
Sum PeCDD	n.d.-10.9	0.5	0.7	n.d.-10.2	0.6	0.4	n.d.-15.1	0.7	3.5*
Sum HxCDD	n.d.-27.5	3.1	0.8	n.d.-15.9	3.8	1.4	n.d.-26.5	3.5	2.3*
Sum HpCDD	0.6-26.2	5.9	0.9	n.d.-18.5	5.9	1.4	n.d.-25.1	6.3	1.5
2,3,7,8-TCDF	n.d.-29.7	0.8	0.2*	n.d.-5.7	1.3	1.3	n.d.-19.9	1.4	0.6
1,2,3,7,8-PeCDF	n.d.-3.3	0.1	0.3*	n.d.-1.1	n.d.	-	n.d.-2.6	0.2	0.5
2,3,4,7,8-PeCDF	n.d.-5.1	0.3	0.4	n.d.-1.7	0.4	1.7*	n.d.-3.4	0.7	0.8
1,2,3,4,7,8-HxCDF	n.d.-3.4	0.3	0.9	n.d.-2.0	0.5	1.4	n.d.-4.8	0.3	1.4
1,2,3,6,7,8-HxCDF	n.d.-4.4	0.1	1.1	n.d.-3.3	0.2	0.7	n.d.-3.4	0.2	1.0
2,3,4,6,7,8-HxCDF	n.d.-4.3	0.3	1.2	n.d.-3.4	0.3	3.6*	n.d.-3.5	0.1	1.1
1,2,3,7,8,9-HxCDF	n.d.-3.5	n.d.	-	n.d.-0.6	n.d.	-	n.d.-0.2	n.d.	0.8
1,2,3,4,6,7,8-HpCDF	0.3-7.4	1.4	1.1	n.d.-7.8	1.7	1.0	0.2-8.4	1.9	1.2
1,2,3,4,7,8,9-HpCDF	n.d.-1.2	n.d.	0.9	n.d.-1.3	0.1	1.0	n.d.-1.4	n.d.	0.6
OCDF	n.d.-6.4	1.0	1.1	n.d.-4.1	0.8	3.5*	n.d.-5.8	0.4	1.0
Sum TCDF	n.d.-134	11.6	0.5	0.5-33.0	12.0	1.3	0.8-78.0	14.8	0.8
Sum PeCDF	n.d.-26.7	3.3	0.5	n.d.-17.5	3.8	1.1	n.d.-19.9	4.6	1.0
Sum HxCDF	n.d.-24.7	2.3	0.9	n.d.-18.2	2.7	0.8	n.d.-19.6	3.7	1.2
Sum HpCDF	0.3-19.6	2.8	1.3	n.d.-11.9	3.6	2.2*	0.5-14.5	3.7	1.3
Sum PCDD	4.7-77.8	22.9	0.9	4.4-164	22.6	1.7*	2.0-109	28.7	1.8*
Sum PCDF	0.3-171	23.6	0.6	3.1-71.9	22.9	1.4	2.0-103	42.2	0.9
Sum PCDD/F	6.0-195	47.2	0.7	7.5-173	45.3	1.1	7.2-181	62.1	1.3
TEQ (I-TEF)	0.02-6.0	0.4	0.5	0.001-2.0	0.7	1.7*	0.04-3.9	0.8	0.8
TEQ (WHO 2005)	0.02-4.9	0.4	0.5	0.002-1.7	0.6	1.7*	0.05-3.1	0.5	0.9

288
 289
 290 Thus, the concentrations observed at three Alpine sites of our study (duration of each sampling period
 291 in 2005-2013: 3 months) were one order of magnitude lower than those monitored in Mediterranean
 292 urban air (Castro-Jimenez et al., 2017; Barbas et al., 2018) and up to one order of magnitude higher than

293 those observed in arctic air at Spitsbergen and at Alert, Canada (duration of each period in winter
294 2000/01: 1 week), where PCDD and PCDF particle-phase concentrations ranged between 2.1 and 13 fg
295 m^{-3} and 2.4 and 46 fg m^{-3} , respectively (Hung et al., 2010). During the four weeks of January 2001,
296 where the highest concentrations at Alert had been monitored, average OCDD and OCDF concentrations
297 were 1.3 and 1.5 fg m^{-3} , respectively. In contrast, OCDD and OCDF multi-annual means (medians) at
298 SBO were 9.36 and 1.1 fg m^{-3} , respectively. It should be mentioned that in the Arctic samples the
299 concentrations of OCDD and OCDF are in the same level, whereas at the Alpine sites the concentrations
300 of OCDD are clearly higher than OCDF concentrations, which fits well to the known congener profile
301 of ambient air data from lowlands. Possibly, the different OCDD/OCDF ratios indicate a pronounced
302 influence of heating activities at the Alpine sites.

303
304 In the winter half years, atmospheric concentrations of some PCDD/F compounds were on average
305 higher than in summer half years (Table 1); this tendency (significant only for some compounds) was
306 observed at the two sites WEI and UFS (which are situated 450 m lower than SBO) but not at SBO.
307 Only at WEI, the winter concentrations, expressed as I-TEQ and WHO-TEQ, were significantly higher
308 than those in summer ($p < 0.05$). The higher the mountain site is, the more it reflects conditions of free
309 atmosphere. Thus, the highest site, SBO is particularly decoupled from the adjacent valleys, especially
310 during winter with low atmospheric convection. Hence, the winter periods are prone to temperature
311 inversions which can trap emissions close to the ground level for several days (Lohmann and Jones,
312 1998). When capping inversions vanish, pollutants can be transported up to higher levels. Additionally,
313 the horizontal distance between SBO to the nearest urban area is greater than in the case of UFS and
314 WEI (Kirchner et al., 2015) and therefore local combustion processes play a minor role at SBO.

315
316 At Austrian low-land sampling sites (Moche and Thanner, 2003), winter/summer ratios of PCDD/F
317 concentrations in the air were in the range of 1.4-7.0; at the mountain site Zöbelboden (900 m a.s.l.), the
318 ratio is 1.6. A connection between cold temperatures and a rise in PCDD/F concentrations in the winter
319 months was found similarly in Baden-Württemberg/Germany (Wallenhorst et al., 1997) and
320 Catalonia/Spain (Parera et al., 2018). In a rural area of Taiwan, the PCDD/F concentration of four
321 seasons from highest to lowest is winter, fall, spring, and summer; ambient air concentration in winter
322 was higher by a factor of 3.4 and 3.9 for total PCDD/F concentration and I-TEQ value (Shih et al., 2006),
323 whereas domestic heating is not found in this region. Around industrial sites in Shanghai, the winter-to-
324 summer ratio of total 2,3,7,8-substituted PCDD/F concentrations was 2.6 due to seasonal variations in
325 sources, environmental processes and meteorology (Die et al., 2015). Compared to these results the
326 winter/summer ratios at the alpine summits of our study are clearly lower.

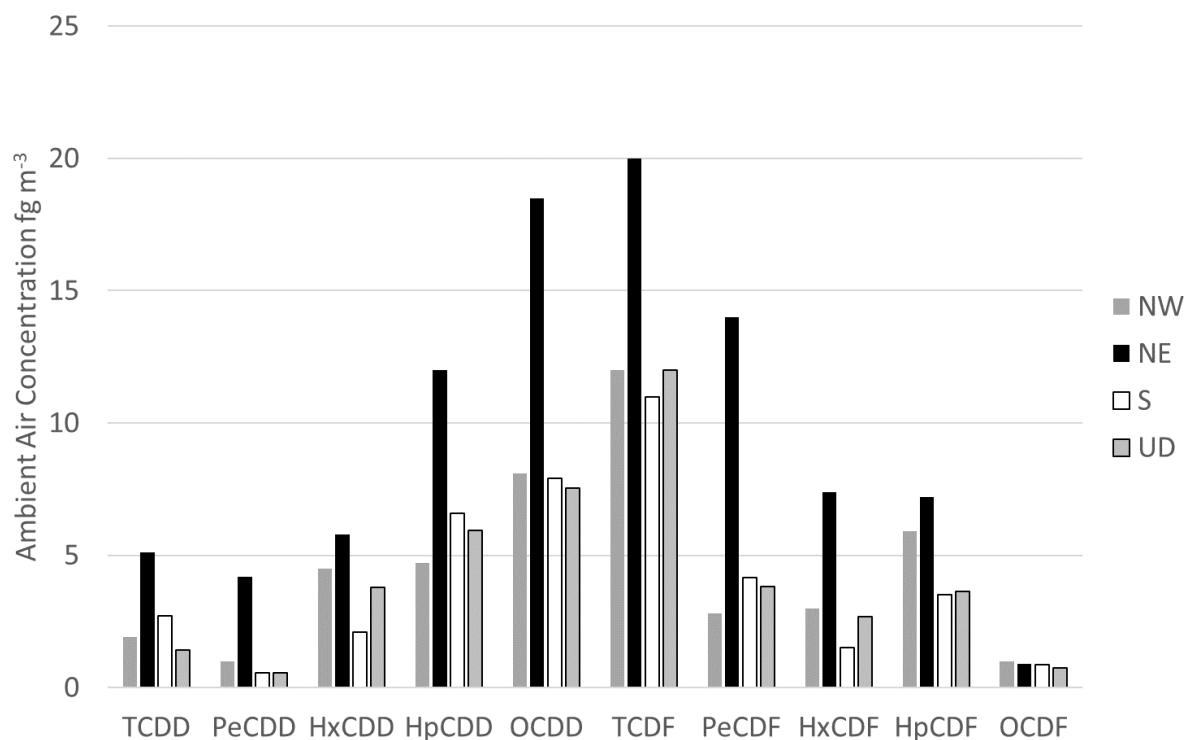
327
328

329 3.3. Median air concentrations in dependency on air masses

330
331 With respect to the time series, there are slight, mostly non-significant concentration differences
332 regarding the four source regions (see Table S1). However, air mass origin was the primary source of
333 the high variability regarding the singular air concentration data, similarly to the results of Sellström et
334 al. (2009).

335
336 For some PCDD/F compounds, concentrations were significantly higher when air masses originated in
337 the NE source region compared with those when air masses originated from elsewhere (NW, S, UD).
338 Transport of PCDD/F to the Alpine region may originate directly from anthropogenic emission areas
339 particularly in eastern central Europe (NE) and indirectly from re-emissions from reservoirs in the soil
340 triggered by temperature-enhanced re-volatilization. At the Swiss station (WEI), higher PCDD/F levels
341 in air masses originating from NE for quite all homologue sums could be determined relatively often
342 (Fig. 2). This may be caused by the influence of sources from the adjacent Klosters-Davos area, western
343 Austria, Bavaria, Czech Republic, and Poland. For OCDD and some congeners or sums of PCDF, the
344 situation is similar at the Bavarian site (UFS), where relatively polluted air is transported via Loisach
345 valley from the Munich area, north-western Austria and eastern central Europe. At the highest site in
346 the Central Alps (SBO), which is situated above less densely populated valleys, PCDD/F concentrations
347 in air coming from NE, NW, and UD are rather balanced. In contrast to WEI and UFS, the sum of
348 PCDD/F measured at SBO does not show any significant dependency from one of four source regions

349 in contrast to WEI and UFS. Median TEQ (I-TEF) is significantly higher at WEI with atmospheric
 350 transport from NE. No differences between the four source regions have been found in the case of SBO
 351 and UFS. However, air masses from the NE source region are significantly less frequent at the three
 352 sites than air masses from the other source regions.
 353



355 Figure 2: Comparison of the PCDD/F homologue pattern of median ambient air concentrations at WEI
 356

357
 358 Air originating from the Mediterranean region was frequently characterized by the lowest concentrations
 359 of PCDD/F. This is in contrast to other POPs, where some organochlorine pesticides (OCPs) are emitted
 360 predominantly in the Mediterranean region and other southern regions: Endosulfan and heptachlor
 361 concentrations show maximum concentrations at UFS, SBO, and WEI with air masses originating from
 362 S (Kirchner et al., 2016). Similarly to PCDD/F and other POPs with (former) industrial and technical
 363 use such as pentachlorobenzene, hexachlorobenzene, and pentachloroanisole have higher concentrations
 364 with air masses from NE.
 365

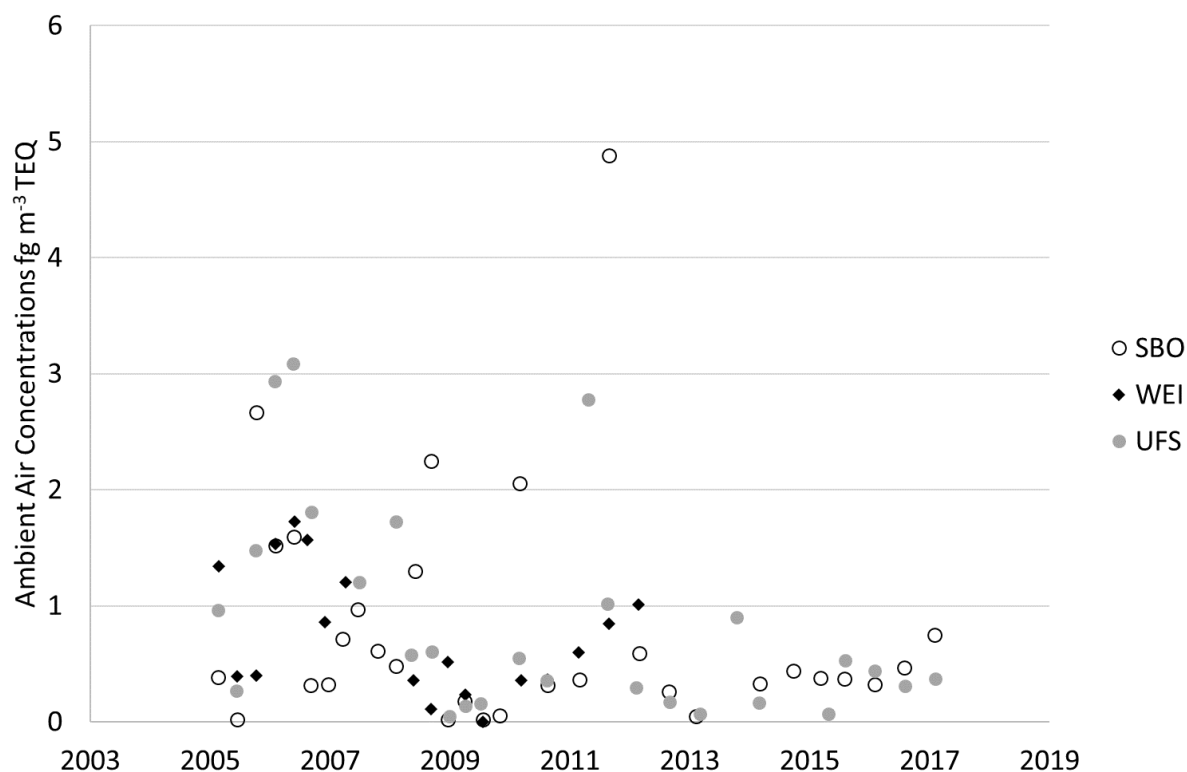
366 Considering the rapid deposition particularly of Penta- to OctaCDD near source areas (Wania and
 367 Mackay, 1996) and the complex long-range transport mechanisms, it is not surprising that differences
 368 between the relatively low concentrations of some PCDD/F congeners, such as 1,2,3,4,7,8-HxCDD,
 369 1,2,3,6,7,8-HxCDD and 1,2,3,7,8,9-HxCDD, for four source regions are relatively small. POPs can be
 370 transported directly or through multiple cycles of atmospheric deposition and subsequent volatilization
 371 from East-Asia via Russia or North America to Europe (Kirchner et al., 2016). For instance, back-
 372 trajectory analysis revealed that relatively high PCDD/F concentrations observed in Alert/Canada in
 373 winter 2000/2001 originated from air masses from Russia/Eurasia (Hung et al., 2002; Sellström et al.,
 374 2009; Hung et al., 2010).
 375

377 3.4. Temporal trend of air concentrations

379 The PCDD/F air concentrations monitored at SBO, WEI and UFS decreased in the first 5 years of our
 380 measurements and increased slightly after 2010 (see Table S2). During 2005-2013, the trend was
 381 negative for some of the selected substances ($p < 0.10$). But only for a few analytes, a significant negative
 382 trend ($p < 0.05$; $p < 0.001$) was identified for this period. Figure 3 shows the temporal trend for the sum

383 of all measured PCDD/F in ambient air given in TEQ. Only for UFS there is a significant decline in the
384 TEQ over the entire period detectable.

385
386 Concerning the single compounds, negative trends were more pronounced at UFS and WEI than at SBO.
387 This may be caused by the different altitudes of three sites and by the fact that global emissions trends
388 play a more important role at SBO as the summit with the highest share of air from the free troposphere.
389 However, after 2013, slightly lower WHO-TEQ concentrations were observed at SBO and UFS in
390 comparison to the first half of the entire dataset.
391



393 Figure 3: Temporal trend of mean weighted concentrations in ambient air for total PCDD and PCDF in
394 WHO-TEQ (2005) for Sonnblick Observatory (SBO) and Schneefernerhaus/Zugspitze (UFS) in 2005-
395 2018, and Weissfluhjoch (WEI) in 2005-2013.

396
397
398 Since the late 1980s, a generally decreasing trend of PCDD/F air concentrations was observed in Europe
399 (Coleman et al., 1997; EPA, 2013). The winter maxima of 847 air samples collected between April 1989
400 and March 1999 at different German sites - without special impact from nearby point sources - decreased
401 by 50 percent (BLAG, 2001; Fiedler, 2003). Time series of PCDD/F in ambient air of a large conurbation
402 in Rhine-Ruhr region (Germany) exhibited a pronounced decrease between 1988 and 2011 (Bruckmann
403 et al., 2013); the concentrations in 1988 ranged from 130 to 332 fg TEQ m⁻³, whereas the levels in 2011
404 ranged from 17 to 24 fg TEQ m⁻³. Bruckmann et al. (2013) stated that the abatement measures at the
405 sources have been successful, also because the differences of air concentrations between the stations
406 had nearly vanished. The strongly decreasing trends monitored in Germany in the 1990s are consistent
407 with those noted for UK urban centers and Catalonia/Spain (Katsoyiannis et al., 2010, Abad et al., 2007)
408 and are caused by considerable emission reductions which may also be a result of the implementation
409 of the POP protocol (EMEP, 2015; van der Gon et al., 2007). According to the information of countries
410 which are part of EMEP (European Monitoring and Evaluation Programme), anthropogenic emissions
411 have decreased from 1990 to 2012 by 60% for PCDD/F. Maximum decrease in annual mean air
412 concentrations of PCDD/F occurred in the beginning of this period, while at the end of this period
413 concentrations levelled off (EMEP, 2015). For Catalonia/Spain a decrease of about 66% - 68% between
414 1994 and 2015 is reported for industrial and traffic sources, whereas slight increases are reported for
415 background sites (Parera et al., 2018). This observation fits very well to our results and indicates that
416 concentrations of PCDD/Fs in ambient air have become more similar between industrial and rural sites

417 in Europe. This supports the conclusion of Dopico and Gómez (2015) that non-industrial sources such
 418 as biomass burning and heating have become relevant sources, since abatement activities have mainly
 419 reduced emissions of industrial sources.

420
 421

422 3.5. Source contributions

423

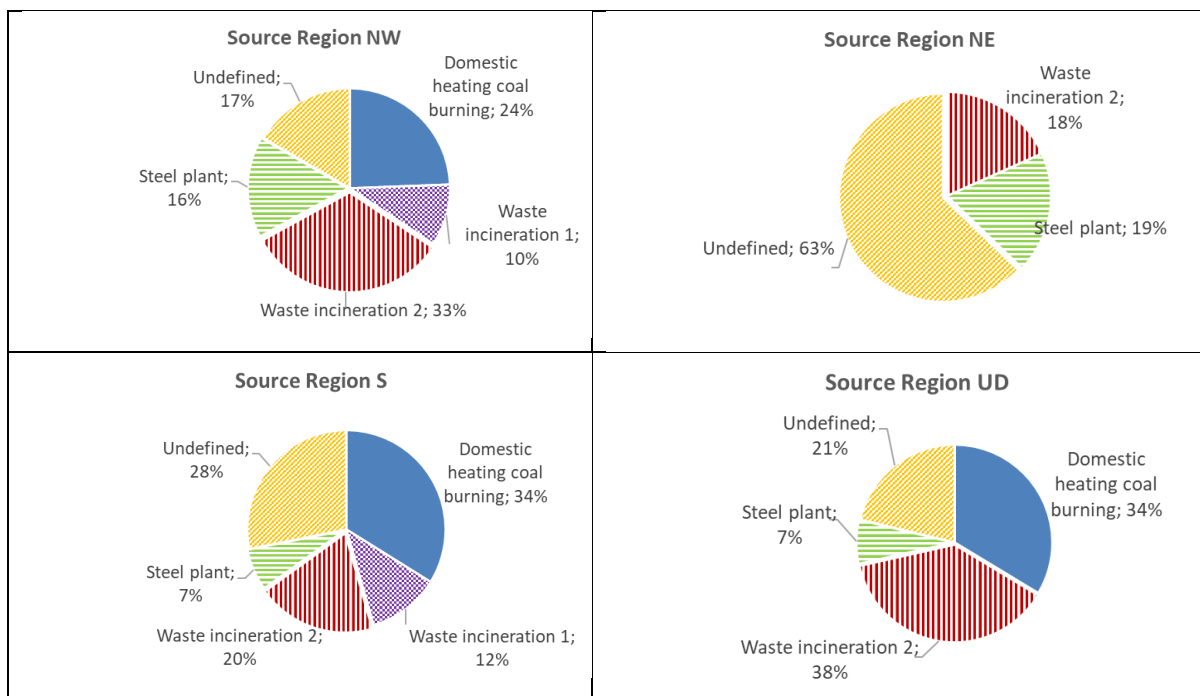
424 In order to estimate source contributions from the observed congener-patterns in ambient air, the EPA
 425 Positive Matrix Factorization 5.0 tool was used (EPA-PMF, USEPA 2014). The four source regions
 426 NW, NE, S, and UD were treated separately, whereas the samples of the three measurement stations
 427 were merged, in order to have a sufficient high amount of data for statistical evaluation. The merging of
 428 data of the three stations is justified because the four source regions are defined for all of them in the
 429 same way. With EPA-PMF the time frame from 2005 till 2015 was evaluated, for which distinct datasets
 430 on the four source regions are given. At minimum, the input-datasets encompassed 17 x 61 data points
 431 (17 congeners and 61 samples). For the equation-based uncertainty the average limit of detection was
 432 used for each congener and an estimated error of ten percent. Missing samples were replaced by the
 433 congener median. Concerning the signal-to-noise-ratio of the statistical approach (S/N) of each
 434 congener, we excluded congeners with S/N smaller than 0.5 by marking them as “bad”. Congeners with
 435 S/N smaller than 1.5 were included to the calculation as “weak”. To execute the PMF in order to
 436 calculate the PMF-factors, we used 20 iterations by using a number of 6 contributing factors, expecting
 437 contributions from waste incineration, steel production, forest fire, coal burning and heating, traffic, and
 438 one undefined source. Congener patterns of the calculated PMF-factors were compared with congener
 439 patterns from the literature to estimate the contribution from different sources (Alcock et al. 1998,
 440 Aristizábal et al. 2011, Castro-Jiménez et al. 2017, Colombo et al. 2009, Colombo et al. 2013, Coutinho
 441 et al. 2015, Denys et al. 2012, Gullet and Touhati 2003, Gunes and Saral 2014, Hagenmaier et al. 1994,
 442 Li et al. 2010, Lin et al. 2007, Mosca et al. 2012, Wang et al. 2013). If no clear pattern could be
 443 distinguished for PMF-factors, the unclear PMF-factors were merged into one which we depicted
 444 “undefined”.

445

446 Up to five contributing sources for PCDD/F in ambient air at the high alpine monitoring stations could
 447 be distinguished by applying the EPA-PMF. Figure 4 depicts the estimated contributions for the
 448 investigated source regions.

449

450



451

452 Figure 4: Source contribution calculated by EPA-PMF for ambient air concentrations of PCDD/Fs.

453
454 As a dominant source of PCDD/F we distinguished domestic heating and coal burning, which was
455 characterized by high concentrations of OCDD, 1,2,3,4,7,8,9-HpCDF and OCDF (Lin et al. 2007, Denys
456 et al. 2012). This source contributed between 24 and 34 percent of PCDD/F ambient air concentrations.

457
458 Another dominant source was waste incineration, where we observed two slightly different patterns:
459 Both patterns of waste incineration are characterized by high concentrations of 1,2,3,6,7,8-HxCDD,
460 1,2,3,7,8,9-HxCDD and 1,2,3,4,6,7,8-HpCDD. However, one waste incineration pattern named “waste
461 incineration 1” did not additionally show high concentrations OCDD which is fitting to a signature for
462 waste incineration reported by Colombo et al. (2009). The other waste incineration pattern “waste
463 incineration 2” includes high concentrations of OCDD which mirrors emissions from municipal waste
464 incineration reported by Wang et al. (2013). Both waste incineration patterns together contributed
465 between 18 and 43 percent of PCDD/F concentrations.

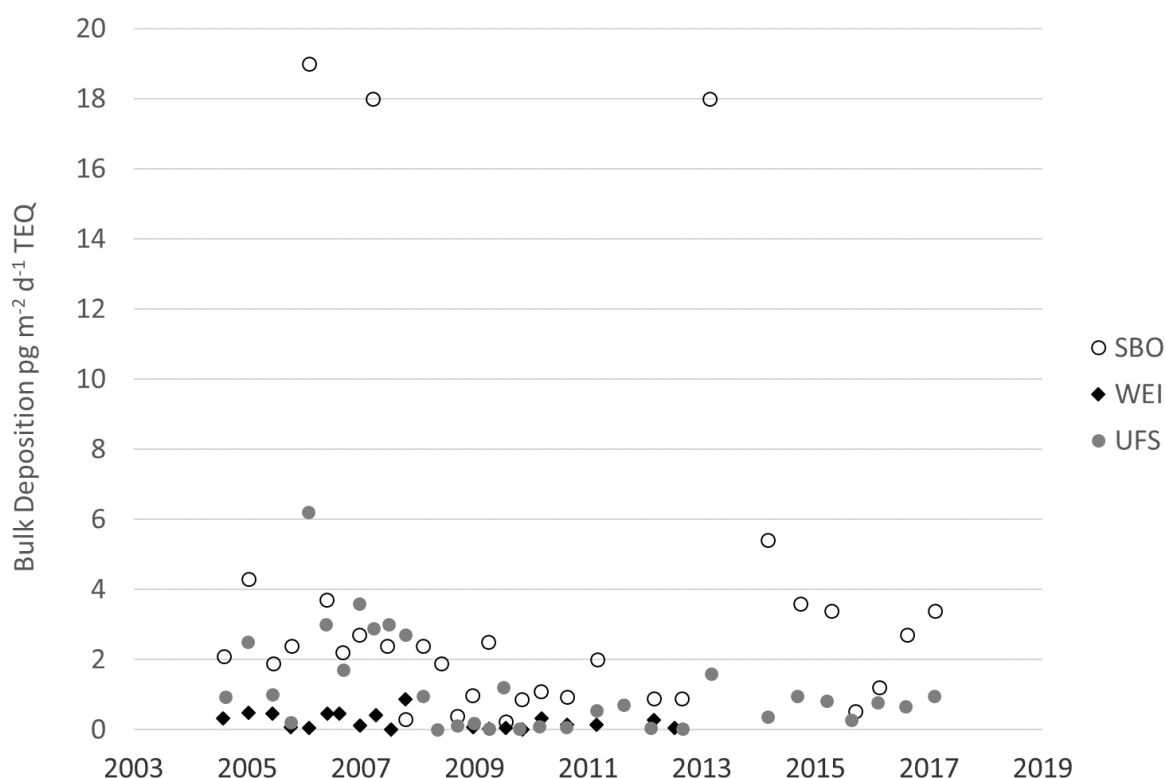
466
467 The third dominant source pattern which was elaborated by the EPA-PMF was characteristic for steel
468 plants, where high concentrations of 2,3,7,8-TCDF, 1,2,3,7,8-PeCDF, and 2,3,4,7,8-PeCDF are
469 indicative (Colombo et al. 2009, Colombo et al. 2013, Coutinho et al. 2015). The emissions from steel
470 plants contributed around 7 percent for the source regions S and UD, and 16 to 19 percent for the source
471 regions NW and NE.

472
473 Using the EPA-PMF we were not able to attribute between 17 and 63 percent of PCDD/F ambient air
474 concentrations to a specific source. These undefined sources did not show explicit characteristics in their
475 congener profiles and may represent long-range atmospheric transport (LRAT) of a mixture of different
476 sources.

477
478 Comparing the four source regions, it shows that the source regions NW, S, and UD show surprisingly
479 similar patterns of source contributions. Only the source region NE is characterized by a huge proportion
480 of 63 percent of PCDD/F ambient air concentrations which cannot be attributed to a specific source.

481 482 3.6. Deposition: Range, seasonality and trend

483
484 Concerning the sum of PCDD/F, medians of bulk deposition rates were significantly different between
485 the three sites ($p < 0.05$) with $321 \text{ pg m}^{-2} \text{ d}^{-1}$ at SBO, $72 \text{ pg m}^{-2} \text{ d}^{-1}$ at UFS, and $30 \text{ pg m}^{-2} \text{ d}^{-1}$ at WEI. The
487 median deposition rates of PCDD/F - calculated in TEQ values (WHO 2005) - were 2.2, 0.8 and 0.14
488 $\text{pg TEQ m}^{-2} \text{ d}^{-1}$, respectively.



492 Figure 5: Bulk deposition rates of PCDD/F in TEQ (WHO 2005) at Sonnblick Observatory (SBO) in
 493 2005 - 2018, Weissfluhjoch (WEI) in 2005 – 2013, and Schneefernerhaus/Zugspitze (UFS) in 2005 –
 494 2018.

495
 496 Using a Spearman rank correlation, deposition fluxes at SBO and UFS show a weak but significant
 497 positive correlation for TEQ ($R=0.503$, $p<0.01$) and Sum PCDD/F ($R=0.395$, $p<0.05$). This indicates
 498 similarities of PCDD/F deposition fluxes at these two sites. However, as established by other studies in
 499 Europe for instance Ispra/Lago Maggiore in Northern /Italy (Castro Jimenez et al., 2012), our results
 500 show a high temporal variability of the PCDD/F atmospheric deposition fluxes (Figure 5).

501
 502 Deposition is dominated in the majority of the samples by the higher chlorinated congeners (see Table
 503 2). The sum of HpCDD, OCDD, HpCDF, and OCDF accounts for 42-97% (Median 74%) of the sum of
 504 PCDD/F. Most dominant is OCDD which on its own accounts for 18-75% (Median 48%) of the sum of
 505 PCDD/F.

506
 507 At SBO, extremely high deposition rates for Sum PCDD/F (4037 and 6439 $\text{pg m}^{-2} \text{d}^{-1}$) were monitored
 508 during two periods (11/2006-3/2007 and 1/2008-4/2008, respectively). These high values do not
 509 coincide neither with high air concentrations at SBO nor with high deposition rates at WEI and UFS.
 510 The pattern of the two samples with extremely high deposition rates are characterized by a high amount
 511 of higher chlorinated congeners (92% and 95%). The most dominant congeners of these two events are
 512 OCDD (with 2800 and 4700 $\text{pg m}^{-2} \text{d}^{-1}$) followed by 1,2,3,4,6,7,8-HpCDD (with 550 and 800 $\text{pg m}^{-2} \text{d}^{-1}$),
 513 OCDF (with 130 and 94 $\text{pg m}^{-2} \text{d}^{-1}$) and 1,2,3,4,6,7,8-HpCDF (with 71 and 68 $\text{pg m}^{-2} \text{d}^{-1}$). This
 514 fingerprint suggests LRAT with no indication of nearby emission sources (Castro-Jimenez et al. 2010,
 515 Castro-Jiménez et al. 2017).

516
 517 The sum of 2,3,7,8-PCDD/F, which is not reported in Table 2, ranged between 10.2 and 5742 $\text{pg m}^{-2} \text{d}^{-1}$
 518 at SBO and is comparable to the deposition flux at Ispra/Lago Maggiore were the total PCDD/F-TEQ
 519 deposition flux monitored during one week was 20 $\text{pg TEQ m}^{-2} \text{d}^{-1}$ in 2005 (Castro-Jiménez et al., 2008).

520
 521 In contrast to air concentrations, the deposition rates determined in our study differ less than 1-2 orders
 522 of magnitude from deposition rates at lowland or urban sites and are partly in the same magnitude as
 523 measurements performed closer to sources. At an urban background site and two rural sites in Bavaria,
 524 one in the sub-alpine region, the medians of PCDD/F deposition rates were in the range of

525 1.7-5.2 pg WHO-TEQ m⁻²d⁻¹ in 2002-2003 (Bayerisches Landesamt für Umwelt, 2006). Urban
526 background deposition rates of PCDD/F of about 10 pg TEQ m⁻² d⁻¹ were found for North-Rhine
527 Westphalia/Germany (Landesanstalt für Umweltschutz Baden-Württemberg, 2004). These deposition
528 rates are almost comparable with the deposition of 5-10 pg WHO-TEQ m⁻²d⁻¹ at the remote site
529 Eifel/Germany (Bruckmann et al., 2013). The total PCDD/F-TEQ deposition flux monitored into Lake
530 Maggiore/Italy during one week in 2005 was 20 pg TEQ m⁻² d⁻¹ (Castro-Jiménez et al., 2008). The bulk
531 deposition of PCDD/F measured at three forest sites in Denmark was in the range of 2.2-3.3 pg TEQ m⁻²
532 d⁻¹; the low variation indicates that PCDD/F have been distributed in the air masses during long-range
533 atmospheric transport (Hovmand et al., 2007). For Japan, deposition rates of 30.1-45.6 and 15.6 pg TEQ
534 m⁻² d⁻¹ were measured in urban and mountain areas, respectively (Ogura et al., 2001). At Lulin/Taiwan,
535 an atmospheric background station (2862 m a.s.l.), the deposition of PCDD/F amounted to 0.7 pg TEQ
536 m⁻² d⁻¹ (Suryani et al., 2015). The total daily dry and wet deposition flux of PCDD/F-TEQ measured in
537 two cities of Northern China in 2014 was in the range of 15-70 pg m⁻² (Zhu et al., 2017).
538
539

540 Table 2: Bulk deposition rates of PCDD/F at Sonnblick Observatory (SBO) and
541 Schneesfernerhaus/Zugspitze (UFS) in 2005-2018, and Weissfluhjoch (WEI) in 2005-2013; Ranges
542 (maximum-minimum), medians and winter to summer ratio calculated on the basis of medians
543 (including outliers)
544

Compound	SBO			WEI			UFS		
	Range pg m ⁻² d ⁻¹	Median pg m ⁻² d ⁻¹	Winter to summer ratio (medians) *p<0.05	Range pg m ⁻² d ⁻¹	Median pg m ⁻² d ⁻¹	Winter to summer ratio (medians) *p<0.05	Range pg m ⁻² d ⁻¹	Median pg m ⁻² d ⁻¹	Winter to summer ratio (medians) *p<0.05
2,3,7,8-TCDD	n.d.-0.4	0.1	-	n.d.-n.d.	n.d.	-	n.d.-0.4	n.d.	-
1,2,3,7,8-PeCDD	n.d.-4.0	0.7	2.3*	n.d.-0.3	n.d.	-	n.d.-1.2	0.2	1.0
1,2,3,4,7,8-HxCDD	n.d.-5.6	0.8	1.5	n.d.-0.5	n.d.	-	n.d.-2.8	0.3	1.5
1,2,3,6,7,8-HxCDD	0.3-16.0	1.7	1.6	n.d.-0.8	n.d.	-	n.d.-7.4	0.6	1.1
1,2,3,7,8,9-HxCDD	0.5-10.0	1.5	1.4	n.d.-0.3	n.d.	-	0.1-5.6	0.8	1.4
1,2,3,4,6,7,8-HpCDD	0.8-800	32.5	1.3	n.d.-14.0	3.3	1.7	1.3-54.0	13.0	1.3
OCDD	3.2-4700	140	0.9	n.d.-n.d.	11.5	1.3	1.5-120	30.5	1.4
Sum TCDD	n.d.-16.0	1.4	2.3*	n.d.-11.0	n.d.	-	n.d.-43.0	1.6	3.9*
Sum PeCDD	n.d.-38.0	3.5	2.2*	n.d.-1.1	n.d.	-	n.d.-29.0	2.0	2.0*
Sum HxCDD	1.7-100	13.5	1.8*	n.d.-3.8	1.0	3.4*	0.5-69.0	7.9	1.3
Sum HpCDD	3.2-1200	54.5	0.8	n.d.-25.0	5.5*	1.4	0.4-83.0	16.0	0.8
2,3,7,8-TCDF	n.d.-5.7	1.1	1.4	n.d.-1.7	0.4	0.5	0.3-8.6	0.6	1.0
1,2,3,7,8-PeCDF	n.d.-5.6	0.7	1.4	n.d.-0.6	n.d.	-	n.d.-1.3	0.3	1.1
2,3,4,7,8-PeCDF	0.2-14.0	1.2	1.4	n.d.-0.6	n.d.	-	n.d.-2.8	0.8	1.4
1,2,3,4,7,8-HxCDF	0.4-22.0	2.1	1.7	n.d.-1.0	0.2	1.5	0.2-6.1	0.7	1.4
1,2,3,6,7,8-HxCDF	0.3-16.0	1.2	1.2	n.d.-1.0	n.d.	-	n.d.-4.5	0.5	1.2
2,3,4,6,7,8-HxCDF	0.1-20.0	1.1	1.4	n.d.-0.7	n.d.	-	n.d.-5.7	0.5	1.5
1,2,3,7,8,9-HxCDF	n.d.-1.8	0.1	1.8*	n.d.-n.d.	n.d.	-	n.d.-0.4	0.1	-
1,2,3,4,6,7,8-HpCDF	1.5-72.0	9.2	1.6	n.d.-3.0	1.2	0.6	0.8-25.0	1.9	1.1
1,2,3,4,7,8,9-HpCDF	0.1-20.0	1.5	1.5	n.d.-0.2	n.d.	-	n.d.-3.1	0.2	2.4*
OCDF	1.2-130	13.0	0.8	n.d.-2.5	0.3	1.0	1.1-9.6	2.5	0.9
Sum TCDF	0.9-83.0	18.0	1.5	n.d.-6.8	2.4	1.1	0.6-38.0	8.7	1.3
Sum PeCDF	1.1-120	13.0	1.5	n.d.-4.7	1.5	2.0	0.6-25.0	6.7	1.4
Sum HxCDF	1.8-150	11.0	1.3	n.d.-2.9	1.3	0.7	0.7-31.0	4.4	1.4
Sum HpCDF	1.5-130	11.0	1.9*	n.d.-5.3	2.3	1.0	0.4-35.0	5.3	1.3
Sum PCDD	8.7-6040	240	0.9	n.d.-110	18.0	1.3	5.0-278	44.0	1.2
Sum PCDF	8.3-466	60.0	1.2	0.6-16.8	8.4	0.9	2.8-117	24.0	1.3
Sum PCDD/F	17.0-6439	321	0.9	1.0-120	30.0	1.2	15.0-369	72.0	1.7*
TEQ (I-TEF)	0.3-22.5	2.5	1.5	n.d.-0.9	0.2	1.7	n.d.-6.2	0.9	1.4
TEQ (WHO 2005)	0.2-19.0	2.2	1.6	n.d.-0.9	0.14	1.7	n.d.-6.2	0.8	1.3

545
546
547 The sum of PCDD/F tends to decrease significantly at UFS and WEI, while WHO-TEQ are declining
548 significantly at UFS only (see Table S3). The deposition rates of the single PCDD/F congeners declined
549 more at UFS than at WEI and SBO.
550

551 The decline of the deposition rates of many PCDD/F compounds is pronounced during the first seven
552 years until 2012. After 2012 we observe a quite constant level or even a slight increase. Particularly at

553 SBO the increase after 2010 is pronounced and there is no trend to lower deposition rates detectable if
554 the whole period from 2005-2018 is considered. For UFS and WEI slightly decreasing trends could be
555 found for the entire period from 2005-2018 and 2005-2015, respectively. Figure 5 shows the temporal
556 trend of deposition of the sum of PCDD/F for the three monitoring stations.

557
558 Within the framework of the European Monitoring and Evaluation Programme (EMEP), PCDD/F
559 deposition decreased more than twice since 1990 (EMEP, 2015). Their analysis indicated that for the
560 Netherlands the reduction of deposition flux from 1990 to 2012 occurred mainly in the beginning of this
561 period and was driven by the reduction of national emissions for PCDD/F. Bruckmann et al. (2013)
562 stated for North-Western Germany, that in parallel to the air concentrations the deposition rates also
563 decreased considerably within the last two decades. However, the decrease of PCDD/F deposition at the
564 remote site Eifel was less pronounced than at sites in urban centers, as important local sources are
565 lacking and the levels are dominated by LRAT. At the Eifel site, the WHO-TEQ deposition revealed a
566 decline by 50%, whereas at the city of Dortmund the deposition of PCDD/F decreased by 75% since
567 1990.

568
569

570 **4. Conclusions**

571
572 The developed air sampling strategy comprising four filter cartridges, which were selected based on
573 trajectory forecast by remote control, worked satisfactory and with a high accuracy with regard to source
574 regions. The sampling campaign showed that air masses with higher PCDD/F concentrations arrived
575 preferably from the northeast region, but due to the lower arrival frequency of air masses from that
576 region their impact on the Alps is of less importance. Thus, none of the predefined source regions is a
577 predominant PCDD/F source for the Alps.

578
579 A clear time trend with decreasing air concentrations could be found from 2005 until 2010, but since
580 2010 the concentrations are increasing again, especially for the most eastern summit (Sonnblick
581 Observatory) where they reach concentrations which are even higher than in 2005. Thus, over the entire
582 sampling period no overarching time trend could be detected. We conclude that the main reduction in
583 PCDD/F ambient air concentrations over Europe happened due to emission reduction measures prior to
584 the start of this monitoring project. The found ambient air concentrations and deposition rates reflect the
585 current situation with nearly unchanged emissions during the last decade.

586
587 Source contribution of measured ambient air concentrations revealed that for the source regions
588 northwest, south, and undefined, waste incineration, heating and coal burning contributed about two
589 thirds of measured concentrations. Only the source region NE showed a distinct pattern where more
590 than half of the measured concentrations could not be assigned to a distinct source and a larger
591 proportion of steel plants played a role.

592
593 The most surprising finding was that deposition rates at the mountainous sites are almost in the same
594 range as in rural lowlands and closer to sources. This was astonishing because the detected ambient air
595 concentrations were one to two orders of magnitude lower than ambient air concentrations found in rural
596 lowlands. A possible explanation for this finding are the higher precipitation rates at the mountainous
597 sites on the one hand and on the other hand the fact that due to the low temperatures the majority of
598 precipitation falls as snow, which has a different scavenging effect than rain. In regard to these findings
599 the question arises if ambient air concentration or deposition measurements give the more accurate
600 picture of the environmental load concerning PCDD/F and POPs in general. Within the Global
601 Monitoring Plan of the Stockholm Convention which has been ratified by all Alpine countries except
602 Italy (Qu et al., 2019) it is foreseen that ambient air concentrations are used to evaluate the POP input
603 into the environment. However, as shown in the present study, ambient air concentrations are not directly
604 proportional to site specific deposition rates. Therefore, it should be discussed if deposition
605 measurements are a necessary component of long-term monitoring in order to assess the environmental
606 burden of POPs, at least at sites with extreme weather conditions such as the Alps or the Arctic.

607

608 To investigate the impact of the measured deposition rates on bioaccumulation of PCDD and PCDF in
609 the food-web of the alpine biosphere, two projects entitled PureAlps have been motivated by the present
610 study. These projects are carried out from 2016 to 2020 by the Bavarian Environment Agency in the
611 surroundings of Mt Zugspitze and by the Environment Agency Austria in the National Park Hohe Tauern
612 in vicinity of the Sonnblick.

613
614

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616
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629 **References**

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