1 Occurrence and source apportionment of perfluoroalkyl acids

(PFAAs) in the atmosphere in China

- 3 Deming Han¹, Yingge Ma², Cheng Huang², Xufeng Zhang¹, Hao Xu¹, Yong Zhou¹, Shan Liang¹,
- 4 Xiaojia Chen¹, Xiqian Huang¹, Haoxiang Liao¹, Shuang Fu¹, Xue Hu¹, Jinping Cheng¹
- ¹ School of Environmental Science and Engineering, Shanghai Jiao Tong University, Shanghai 200240, China
- 6 ² State Environmental Protection Key Laboratory of the Formation and Prevention of Urban Air Pollution Complex,
- 7 Shanghai Academy of Environmental Sciences, Shanghai 200233, China
- *Correspondence to*: Jinping Cheng (jpcheng@sjtu.edu.cn)

Abstract:

Perfluoroalkyl acids (PFAAs) are a form of toxic pollutant that can be transported across the globe and accumulated in the bodies of wildlife and humans. A nationwide geographical investigation considering atmospheric PFAAs_via XAD_Passive Air Sampler was conducted in 23 different provinces/municipalities/autonomous regions in China, which provides an excellent chance to investigate their occurrences, spatial trends, and potential sources. The total_atmospheric concentrations of thirteen PFAAs_(n=268) were 6.19–292.57 pg/m³, with an average value of 39.84±28.08 pg/m³, which were higher than other urban levels but lower than point source measurements. Perfluorooctanoic acid (PFOA) was the dominant PFAAs (20.6%), followed by perfluorohexanoic acid (PFHxA), perfluorooctane sulfonate (PFOS), and perfluoroheptanoic acid (PFPeA). An increasing seasonal trend of PFAAs concentrations was shown as summer < autumn < spring < winter, which may be initiated by stagnant meteorological conditions. Spatially, the content of PFAAs displayed a declining gradient trend of central China> northern China> eastern China> northeast of China> southwest of China> southern China areas, and Henan contributed as the largest proportion of PFAAs. Four sources of PFAAs were identified using a positive matrix factorization (PMF) model, including PFOS-based products (26.1%), PFOA-based, and PFNA-based products (36.6%), degradation products of fluorotelomere-based products (15.5%), and an unknown source (21.8%).

1.Introduction

Perfluoroalkyl acids (PFAAs) are one class of ionic polyfluoroalkyl substances (PFASs), which have excellent characteristics in terms of chemical and thermal stability, high surface activity, and water and oil repulsion (Lindstrom et al., 2011; Wang et al., 2014). They are applied to a wide variety of domestic and industrial products such as textiles, oil

and liquid repellents, firefighting foam, pesticides, and food packaging materials (Xie et al., 2013; Wang et al., 2014). PFAAs can be released to the surrounding environment during manufacturing and use of PFAAs containing products, which are ubiquitous in the environment (e.g., in the atmosphere, water, or snow) (Dreyer et al., 2009; Wang et al., 2017; Hu et al., 2016), in wildlife (Sedlak et al., 2017), and even in the human body (Cardenas et al., 2017; Tian et al., 2018). PFAAs can change adult thyroid hormone levels, reduce newborn birth weight, and biomagnify up the food chain, which can be extremely toxic to animals and humans (Hu et al., 2016; Jian et al., 2017; Baard Ingegerdsson et al., 2010). Of the PFAAs, the long-chain $(C \ge 8)$ perfluoroalkyl carboxylic acids (PFCAs) and $(C \ge 7)$ perfluoroalkyl sulfonic acids (PFSAs) are more toxic and bio-accumulative than their short-chain analogues (Buck et al., 2011). This especially applies to perfluorooctanoic acid (PFOA), perfluorooctane sulfonate (PFOS) and perfluorohexane sulfonate (PFHxS), in which PFOS and PFOA have been added to Annex B and Annex A of the Stockholm Convention in 2009 and 2019, respectively, while PFHxS was under review by the Persistent Organic Pollutants Review Committee (Johansson et al., 2008; UNEP Stockholm Convention, 2019). PFAAs can originate from direct sources of products' emissions as well as indirect sources of incomplete degradation of their precursors. It is estimated that the global historical emission quantities of C4-C14 PFCAs were 2610-21400 t in the period of 1951-2015, of which PFOA-based and perfluorononanoic-acid (PFNA)-based products contributed the most (Wang et al., 2014). A trend of geographical distribution of major fluorochemical manufacturing sites has shifted from Western Europe, US, and Japan to the emerging economies in the Asia Pacific area over the past decades. This is especially true for China, which was the world's largest industrial contributor of PFOAs (50-80 t) and PFOS-related compounds (~1800 t) in 2009 (Xie et al., 2013). PFOA- and PFOS- based products were added to the Catalogue for the Guidance of Industrial Structure Adjustment in China in 2011, and restricted elimination of PFOA/PFOS substances production were conducted. With a large quantity of PFAAs and their products manufacturing and consumption, China has become the emerging contamination hotspots in the world. In spite of several studies on atmospheric PFAAs levels having been conducted in a few cities (Liu et al., 2015) and point sources (Yao et al., 2016a; Tian et al., 2018) in China, due to the imbalanced urbanization and industrialization levels, there is still a lack of systemic research on atmospheric PFAAs quantification and trends in China. Additionally, the long range or mesoscale transport was also suggested to have a contribution to PFAAs in the air (Dreyer et al., 2009; Cai et al., 2012a). In general, three pathways/hypotheses for the transportation of PFAAs were suggested: transport associated with particles, degradation from precursor, and sea salts from current bursting in coastal areas. The PFAAs precursors such as fluorotelomere alcohols (FTOHs), which can form the corresponding PFAAs through oxidation reactions initiated by hydroxyl radicals (OH·) in the atmosphere (Thackray and Selin, 2017), are more volatile

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

than PFAAs and can reach remote areas via long-range transportation (Martin et al., 2006; Wang et al., 2018). Due to the lower acid dissociation coefficient (p K_A), 0-3.8 for PFCAs and -3.3 for PFSAs, PFAAs are expected to be mainly associated with aerosols in the non-volatile anionic form (Lai et al., 2018; Karásková et al., 2018). However, recent field studies have confirmed their occurrence in gaseous phase (Cassandra et al., 2018; Ahrens et al., 2013), e.g. Fang et al., (2018) found the total concentrations of C2, C4-C10 PFCAs and C6 and C8 PFSAs in the gas phase were 0.076-4.0 pg/m³ in the air above the Bohai and Yellow Seas, China. Investigating the transport pathways of PFAAs in nationwide region via active air sampler (AAS) is challenging, due to their electronic power supply and high cost. Fortunately, a number of reports showed that the XAD (a styrene-divinylbenzene copolymer) impregnated sorbent based passive air sampler (SIP-PAS) and XAD based PAS (XAD-PAS), were proven to be an ideal alternative sampling tool for monitoring PFAAs in a wide region. Despite several publications suggested XAD-PAS collects primarily gaseous PFAAs (Melymuk et al., 2014; Lai et al., 2018) in the ambient, current findings were not consistent. Due to the unimpeded movements of particles into the sampler, XAD-PAS was indicated to collect a representative sample of both gas and particle phases (Ahrens et al., 2013; Okeme et al., 2016; Karásková et al., 2018). Moreover, the dominant sorbent for fluorinated compounds was reported as XAD resin in the XAD impregnated SIP-PAS, instead of PUF themselves (Krogseth et al., 2013). XAD-PAS give PFASs profiles that were more closely resembled to those from AAS in comparing with PUF-PAS, have sufficient uptake rates for the PFCAs and PFSAs to be deployed for short time duration (Lai et al., 2018). Given the factors mentioned above, we conducted a nationwide survey of PFAAs in China at a provincial level using a XAD-PAS from January to December in 2017. The objective of this research was: (1) to examine the tempo-spatial variations of PFAAs, and (2) to identify their potential affecting factors and evaluate the affecting pathways. To the best of our knowledge, this is the first research paper analyzing both a long-term and nationwide atmospheric PFAAs data set complemented by a comprehensive investigation in China.

2.Material and methods

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

2.1 Chemicals and reagents

The PFAAs standards used were Wellington Laboratories (Guelph, ON, Canada) PFAC–MXB standard materials, including C5–C14 PFCAs analogues (Perfluoropentanoic acid (PFPA), Perfluorohexanoic acid (PFHxA), Perfluoroheptanoic acid (PFHpA), PFOA, Perfluorononanoic acid (PFNA), Perfluorodecanoic acid (PFDA), Perfluoroundecanoic acid (PFUdA), Perfluorododecanoic acid (PFDA), and

Perfluorotetradecanoic acid (PFTeDA)), as well as C4, C6, and C8 PFSAs analogues (Perfluorobutane sulfonic acid (PFBS), PFHxS, and PFOS). The mass-labeled 1,2-13C2-PFHxA, 1,2,3,4-13C4-PFOA, 1,2,3,4,5-13C5-PFNA, 1,2-13C2-PFDA, 1,2-13C2-PFUdA, 1,2-13C2-PFDOA, 18O2-PFHxS, and 1,2,3,4-13C4-PFOS were used as internal standards (ISs, MPFAC-MXA, Wellington Laboratories Inc.) in high-performance liquid chromatography (HPLC) coupled with a tandem mass spectrometer (MS/MS). HPLC-grade reagents that were used include methanol, ethyl acetate, ammonia acetate, acetone, methylene dichloride, n-hexane, and Milli-Q water. Detailed sources of the target PFAAs and their ISs are listed in Table S1 in the Supplementary Materials.

2.2 Sample collection

87

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

Sampling campaigns were carried out at 23 different provinces/municipalities/autonomous regions in China simultaneously from January to December 2017, of which 20 were urban sites and three were rural sites (Zhejiang, Shanxi, and Liaoning). Urban samples typically came from urban residential areas, and the rural samples were obtained from villages. These sampling sites were divided into seven administrative divisions: norther China (NC, n=3 sites), southern China (SC, n=2), central China (CC, n=3), eastern China (EC, n=7), northwest of China (NW, n=3), northeast of China (NE, n=2), and southwest of China (SW, n=3). A geographical map of the sampling sites is displayed in Figure S1, and the detailed information on sampling sites such as elevation, meteorological parameters, local resident population and gross domestic product were listed in Table S2 and Figure S1. Samples were collected with Amberlite XAD-2 resin using XAD-PAS, which have been successfully monitored PFCAs (C4-C16) and PFSAs (C4-C10) in the atmosphere (Krogseth et al., 2013; Armitage et al., 2013). Briefly, the mesh cylinder (L.× I.D.: 10 cm × 2 cm) was prebaked at 450°C for 3 h, filled with ~10 g XAD-2 resin, and capped with an aluminum cap. The particle size of XAD-2 is ~20-60 mesh, with water content of 20%-45%, its specific surface area \geq 430 m²/g, and the reference adsorption capacity \geq 35 mg/g. We should keep in mind that the unimpeded movement of particle bound PFAAs would be captured during sampling using XAD-PAS, which cannot differentiate PFAAs between gas and particle phases. Despite some research suggest the sampling efficiency of gas and particle phase PFAAs were similar (Karásková et al., 2018). In the present study, the two phases PFAAs sampled by XAD-PAS were treated as the whole atmosphere PFAAs concentration. The sampling program for each sample lasted approximately a month (30 days), and the error of the sampling time was controlled within 3 d. At the end of each deployment period, the atmosphere samples were retrieved, resealed in their original solvent-cleaned aluminum tins at the sampling location, and transported by express post to Shanghai Jiao Tong University. On receipt, they were stored and frozen (-20 °C) until extraction.

The sampling rate of XAD-PAS is a crucial factor to derive the chemical concentrations accumulated in the XAD resin.

Ahrens et al. (2013) found that sampling rate of PFCAs and PFASs ranged 1.80–5.50 m³/d with XAD impregnated sorbent, and the sampling rate increased as the carbon chain adding, while Karásková et al. (2018) suggested that the sampling rate of XAD–PAS of 0.21–15.00 m³/d for PFAAs. The loss of depuration compounds could be used to calculate the sampling rate, assessing the impacts from meteorological factors like temperature and wind speed. According to Ahrens et al. (2013) the 1,2,3,4–1³C₄–PFOA was used to calculate the sampling rates of PFAAs at Shanghai sampling site (Shanghai Jiao Tong University) in the present study, by assessing 1,2,3,4–1³C₄–PFOA abundance loss. The specific description of the sampling rate calculation in this study is shown in Section S1 in the Supplementary Materials.

2.3 Sample preparation and instrument analysis

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

The sample preparation and analysis were according to the method described by previous researches (Liu et al., 2015; Tian et al., 2018). The MPFAC-MXA ISs mixture surrogates (10 ng) were added to each spiked sample prior to extraction. This was done to account for the loss of substances from the samples associated with instrument instability caused by the changes in laboratory environmental conditions. The XAD resin samples were Soxhlet-extracted for 24 h using a Soxhlet extraction system, with n-hexane: acetone (1:1, V:V) as a solvent in a 300 mL polypropylene (PP) bottle, following extracted with methanol for 4 h. These two extracts were combined and reduced to ~5 mL via a rotary evaporator (RE-52AA, Yarong Biochemical Instrument Inc., Shanghai, China) at a temperature below 35 °C, and then transferred to a 10 mL PP tube for centrifugation (10 min, 8,000 rpm). The supernatant was transferred to another PP tube, filtered three times through a 0.22 µm nylon filter, with an addition of 1 mL methanol each time. The extracts were further condensed under a gentle stream of nitrogen (99.999%, Shanghai Liquid Gas Cor.) at 35 °C to a final 200 μL for instrument analysis. The separation and detection of PFAAs were performed using a HPLC system (Thermo Ultra 3000⁺, Thermo Scientific, USA) coupled with a triple quadrupole negative electrospray ionization MS/MS (Thermo API 3000, Thermo Scientific, USA). An Agilent Eclipse XDB C18 (3.5 μm, 2.1 mm, 150 mm) was used to separate the desorbed substances. The column temperature was set to 40 °C, and the flow rate was 0.30 mL/min. The injection volume was 20 μL. The gradient elution program of the mobile phase A (5 mmol/L aqueous ammonium acetate) and B (methanol) was 80% A + 20% B at the start, 5% A + 95% B at 8 min, 100% a at 13 min, 80% A + 20% B at 14 min, and was maintained for 6 min. The MS/MS was operated in a negative ion scan and multiple reaction monitoring (MRM) mode, and the electrospray voltage was set to 4500 V. The ion source temperature was 450 °C. The flow rates of the atomization gas and air curtain gas was 10 and 9 L/min, respectively. Species identification was achieved by comparing the mass spectra and retention time of the chromatographic peaks with the corresponding authentic standards.

2.4 Quality assurance and quality control

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

162

163

164

165

166

167

168

169

170

171

172

173

To avoid exogenous contamination, the XAD-2 resin was precleaned using a Soxhlet extraction system with acetone and petroleum ether at extraction times of 24 h and 4 h, respectively. The extracted XAD resin was dried under a vacuum desiccator, wrapped in an aluminum foil and zip-lock bags, and stored at -20 °C to avoid contamination. All laboratory vessels were PP, and these vessels were washed with ultrapure water and methanol three times, respectively. For quantification, six-point calibration curves of PFAAs were constructed by adopting different calibration solutions with values of 1, 3, 6, 15, 30, and 60 ng/mL. The same concentration for the internal calibration (10 ng/mL) was used for each level of the calibration solution. Recovery standards were added to each of the samples to monitor procedural performance, and the mean spiked PFAAs recoveries ranged from 81%±25% to 108%±22%. All the analyzed PFAAs were normalized against the recovery of the corresponding mass-labeled ISs. Field blanks were prepared at all sampling sites, transported, and analyzed in the same way as the samples. Laboratory blanks were obtained by taking amounts of solvent via extraction, cleanup, and analysis. A total of 8 field blanks and 26 laboratory blanks were analyzed, with individual blank values of BDL (below detection limit)-1.12 pg/m³ and BDL-1.29 pg/m³, respectively. All the results were corrected according to the blank and recovery results. The method detection limit (MDL) was derived from three times standard deviation of the field blank values. The limit of detection (LOD) and the limit of quantification (LOQ) were determined as a signal-to-noise ratio of 3:1 and 10:1, respectively (Rauert et al., 2018;Liu et al., 2015). To convert MDLs, LODs and LOQs values to pg/m³, the mean volume of sampling air (m³) was applied. For the analytes that were not detected or were below the LOQs in field blanks, MDLs were derived directly from three times the corresponding LODs. More detailed information on the individual compounds of PFASs on MDL, LOD, LOQ, the recovery values, and blank values are listed in Table S3.

2.5 Statistical and geostatistical analysis

Statistical analyses were carried out by SPSS Statistics 22 (IBM Inc. US), and the values of 1/2 MDL were used to replace these measured results of BDL. The statistics figures were depicted using technical software of SigmaPlot 14.0 (Systat Software, US). And the geographical variations of atmospheric PFAAs were analyzed with ArcGIS 10.4 (ESRI, US). The Hybrid Single-Particle Lagrangian Integrated Trajectory (Hysplit) back trajectory model (NOAA, US) was used to study the long range transport of air masses in the sampling locations (Zhen et al., 2014). Positive matrix factorization (PMF) is considered an advanced algorithm among various receptor models, which has been successfully applied for source identification of environmental pollutants (Han et al., 2018;Han et al., 2019). PMF (5.0, US EPA) was adopted to cluster the PFAAs with similar behaviors to identify potential sources, and a more detailed description of PMF can be

seen in Section S2.

174

175

176

3. Results and discussion

3.1 Abundances and compositions

177 The descriptive statistics of all targeted atmosphere PFAAs (n=268) are presented in Table 1 and Table S4. The total concentrations of Σ_{13} PFAAs analogues varied between 6.19 and 292.57 pg/m³, with an average value of 39.84±28.08 178 179 pg/m³. The commonly concerned PFCAs analogues (C5-C14) occupied 79.6% of the total PFAAs, at a level of 180 4.50–247.23 pg/m³, whereas the PFSAs concentrations were 1.04–42.61 pg/m³. The long-chain PFCAs concentrations were 17.96±13.71 pg/m³, which were significantly higher than the short-chain concentrations (13.74±12.19 pg/m³) 181 182 (p<0.05). Similarly, a recent PFAAs measurement conducted in the landfill atmosphere in Tianjin, China (Tian et al., 2018), found the long chain PFCAs were much higher than the short species. Specifically, PFOA was the dominant 183 184 PFAAs (accounting 20.6%), and was detected in all atmospheric samples with an average value of 8.19±8.03 pg/m³. This phenomenon could occur since PFOA is widely used in the manufacturing of polytetrafluoroethylene (PTFE), 185 186 perfluorinated ethylene propolymer (FEP), and perfluoroalkoxy polymers (PFA) (Wang et al., 2014). The domestic 187 demand for and the industrial production of PFOA-based products have been increasing in China since the late 1990s 188 (Wang et al., 2014), and direct emissions of FOSA-based products may contribute to the relative high level of PFOA. 189 Meanwhile, one major variation of PFOA precursor, 8:2 FTOH, was reported to rank as the highest concentration among 190 neural PFASs in air of China (De Silva, 2004; Martin et al., 2006). Among PFAAs' composition profile, it was followed 191 by PFHxA, PFOS, and PFPeA, with mean concentrations of 5.36, 5.20, and 4.95 pg/m³, respectively. The detection 192 frequencies of PFCAs decreased gradually as the carbon chain length increased - for instance, the PFPeA and PFTrDA 193 were detected in 84.8% and 37.3%, respectively. 194 Compared with other gaseous PFAAs measurements, Liu et al. (2015) reported that PFAAs in the urban atmosphere 195 sampled with XAD-containing sorbent in Shenzhen city in China was 15±8.8 pg/m³, which contributed to nearly half of 196 this study. Wong et al. (2018) found that a much lower PFAAs levels in the remote Arctic area than this study, with mean 197 value of 1.95 pg/m³. This study found generally higher PFAAs abundances compared to measurement in Canada 198 (Gewurtz et al., 2013), which may be attributed to the relative high abundance of industrial and domestic emissions in 199 China. However, the PFAAs concentrations in urban/rural areas in this study were far lower than the measurements at 200 point sources, for example, landfill atmosphere (Tian et al., 2018) (360–820 pg/m³) and fluorochemical manufacturing 201 facility (Chen et al., 2018) (4900±4200 pg/m³), suggesting that PFAAs were susceptible to being affected by local source

emissions. Although there existed inherent differences of PFAAs levels between regions, the impacts from differences in sampling techniques and sorbents between XAD-PAS and SIP-PAS could not be neglected. As indicated by previous researches, XAD has much higher sorptive capacity of PFASs than PUF, wind speed and temperature displayed different degrees of impact on their sampling capacity among different regions. Additionally, UV radiation has the potential to degradate PFAAs due to O₃, OH·, and other atmospheric oxidants during sampling.

Table 1. Comparison of PFAAs levels in the present research with measurements in other areas (pg/m³)

Sampling sites	Duration	Sampling location	Sampler type ^a	PFAAs ^b	PFCAs ^c	Reference
23 provinces in	2017.1–12	Urban and rural areas	XAD-PAS	6.19–292.57;	4.50–247.23;	This study
China				39.84±28.08	31.69±23.88; C5–C14	
Shenzhen, China	2011.9–11	Urban area	SIP-PAS	3.4–34; 15±8.8	11.59±8.74; C4–C12	(Liu et al., 2015)
Fuxin, China	2016.9–10	Fluorochemical	SIP-PAS	4900±4200	4900±4200; C4–C12	(Chen et al., 2018)
		manufacturing facilities				
Tianjin, China	2013	Waste water treatment	SIP-PAS	87.9–227; 123	87.9–227; 123; C6–C12	(Yao et al., 2016a)
		plant				
Tianjin, China	2016.5-6	Landfill	SIP-PAS	280-820	280-820; C4-C12	(Tian et al., 2018)
Canada	2006–2011	Remote and urban areas	SIP-PAS	0.014-0.44	0.014-0.44; C8-C12	(Gewurtz et al.,
						2013)
Alert, Arctic	2006.8–2015	Remote area	SIP-PAS	1.95	1.95; C4–C8	(Wong et al., 2018)
	.2					
Toronto, Canada	2010.3–10	Semi-urban site	SIP-PAS	11.24±7.95	<u>11.24±7.95;</u> C4–C18	(Ahrens et al., 2013)
Brno, Czech	2013.4-9	Suburban background	XAD-PAS	30–153	26–147.6; C4–C14	(Karásková et al.,
Republic		site				<u>2018)</u>

^a: SIP-PAS represent XAD impregnated PUF sorbent based PAS, which is composed of PUF, mashed XAD-4, and PUF;

3.2 Temporal variations

Monthly and seasonal variations of the mean PFAAs concentrations are depicted in Figure 1. In general, an increasing seasonal mean of PFAAs concentrations from 23 sampling sites existed for summer (31.35 pg/m³) < autumn (35.63)

b: represent the total concentration ranges of PFCAs and PFSAs; mean concentrations of the total PFCAs and PFSAs;

c: represent concentration range; mean value; carbon length of PFCAs.

pg/m³) < spring (42.40 pg/m³) < winter (52.83 pg/m³). The winter maxima abundance of PFAAs could be attribute to the stagnant atmospheric conditions, in which atmospheric contaminants were trapped in the air with a weak diluting effect. XAD-PAS showed similar efficiency of capturing gas and particle phases PFASs, while the unimpeded particle gathering efficiency is challenging to quantify. In addition, despite the increase in atmospheric oxidation of precursors in summer may lead to PFCAs rise (Li et al., 2011; Yao et al., 2016a), the abundant rainfall would enhance their scavenging activities (Table S5), ultimately leading to the relatively low concentrations of PFAAs in the summer. Specifically, the PFAAs showed much higher concentrations in spring than other seasons in Shanghai, which was different from Tianjin and Xinjiang (Figure S2). An extreme high level of PFAAs of 135.51 pg/m³ was occurred in November in Beijing, which was 2–4.5 times higher than in other month, indicating the potential point source of PFAAs contamination in this site. In fact, numerous fluoride related products manufacturers were distributed in EC, NC (including Beijing) and CC areas, see detail in Figure S3. As gaseous PFAAs measurements were majorly reported at a relative short time (several weeks to several months), it is somewhat difficult to compare their temporal trends. Interestingly, the evolution of PFAAs showed a dramatic monthly variation, and the monthly mean levels varied from 25.92 to 60.57 pg/m³, with the lowest and the highest abundances being present in September and December, respectively. For the specific composition profile of PFAAs, the average concentrations of PFOA, PFHxA, PFPeA, and PFOS were 10.36, 8.42, 6.55, and 6.44 pg/m³ in winter, respectively, which were nearly two times higher than in the summer. The seasonal variation trend of PFOS was summer < spring ≈ autumn < winter, while PFNA appeared to show winter maxima with concentrations 4 and 3 times higher than in the summer and spring, respectively. However, Wong et al. (2018) reported that PFBS showed the maximal value in winter but found no consistent seasonality for PFOS in the Arctic area. The differences may be explained as the PFAAs in air in the remote Arctic area were originated from long-range transport and volatilization from snow or sea, but not affected by local direct anthropogenic emission.

215

216

217

218

219

220

221

222

223

224

225

226

227

228

229

230

231

232

233

234

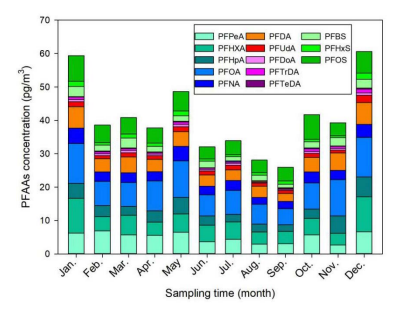


Fig. 1. Monthly mean concentrations of PFAAs in China from January to December 2017

3.3 Geographical distributions

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251

252

253

254

255

Due to the stark differences in topography and socioeconomic development of Chinese provinces, municipalities, or autonomous regions, as well as the enormous differences in industrialization and emissions, PFAAs showed significantly different distribution patterns in China (Figure 2). Overall, the predominant declining gradient of PFAAs' contents was CC> NC> EC> NE> SW> NW> SC areas in China, which was similar to previous research that the outdoor dust-bounded PFAAs were relatively enriched in the eastern part of mainland China (Yao et al., 2016b). This trend was not surprising since numerous PFAAs related photoelectric industries, chemical industries, and mechanical industries are dispersed across CC, EC and NC areas, e.g., Shanghai, Zhejiang, Fujian, Henan, and Jiangsu. As expected, the western mountain and highland areas, e.g., Xinjiang and Yunnan (20.88 pg/m³), with relatively low population densities and high latitudes, displayed significantly lower PFAAs concentrations. It was reported that high orographic conditions have a cold trapping effect on atmospheric PFASs, the transportation of PFAAs involving particles or not should be dramatically reduced (Konstantinos et al., 2010; Yao et al., 2016a). Given that altitudes increase gradually from several meters in EC, NC and SC coastal areas to nearly 2,000 meters in SW and NW highland regions in China, the high altitude blocking effect for atmospheric PFAAs transportation should not be neglected. The annual average concentrations of PFAAs at the provincial level ranged from 12.38 pg/m³ in Xinjiang to 90.88 pg/m³ in Henan, and the composition patterns varied widely. Henan contributed the largest proportion of PFAAs in China, and showed the highest PFOA level (19.07 pg/m³), which is a typical, heavily-industrialized province characterized by textile

treatments, metal plating, and firefighting foam manufacturing, and a large amount of PFAAs emulsifier fluoropolymers were used in industrial production. Special attention should be paid to Zhejiang, the level of which (61.68 pg/m³) ranked second in PFAAs abundances in spite of its sampling site being located in a village. As well as this, several painting-packaging plants, mechanical plants, and electrical equipment manufacturers were dispersed around this sampling site (see Figure S4), which would contribute to the PFAAs variations in this site. In fact, the GDP of Zhejiang ranked fourth in China, specializing in mechanical manufacture, textiles, and chemical industry. Moreover, the top six sites with abundant of PFAAs were located in the most economically-developed and populated areas (the Yangtze River Delta area, the Circum-Bohai Sea Region), and in the rapidly-developing regions (Henan, Sichuan) in China. In line with this result, a sampling campaign conducted in Asia, including 18 sites in China, found very high levels of PFAAs precursors (8:2 FTOH, 10:2 FTOH) existed in Beijing, Tianjin, and Zhejiang (Li et al., 2011). But meanwhile we should keep in mind that the production of PFCAs in the atmosphere from gaseous precursors degradation may be impaired in urban areas, due to the high abundance of NOx compete for OH· radicals. Furthermore, PFOA concentrations were apparently high in Henan, Zhejiang, Beijing, Tianjin, and Hubei, where mean values ranged of 11.65–19.14 pg/m³ compared with in other provinces (2.93–8.54 pg/m³). PFOA and PFOA-related products have not been banned for use in various industrial and domestic applications (Konstantinos et al., 2010; Wang et al., 2014), which were manufactured extensively in EC and NC areas and were used widely. However, the highest concentration of PFOS was found in Zhejiang (14.13 pg/m³), which may be affected by local manufacturing of PFOS based products, e.g. leather, paper and metal plating. It was followed by Beijing (8.98 pg/m³) and Fujian (9.09 pg/m³), while Xinjiang and Yunnan shared the lowest levels (1.20–3.57 pg/m³). This spatial variation patterns of PFOS in the present study, matched well with a previous national survey that found most PFOS and its derivative facilities in China are suited in EC, CC and NC areas, with emission density ranged from 1-500 g/(km²·a) (Konstantinos et al., 2010; Wang et al., 2014).

256

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

272

273

274

275

276

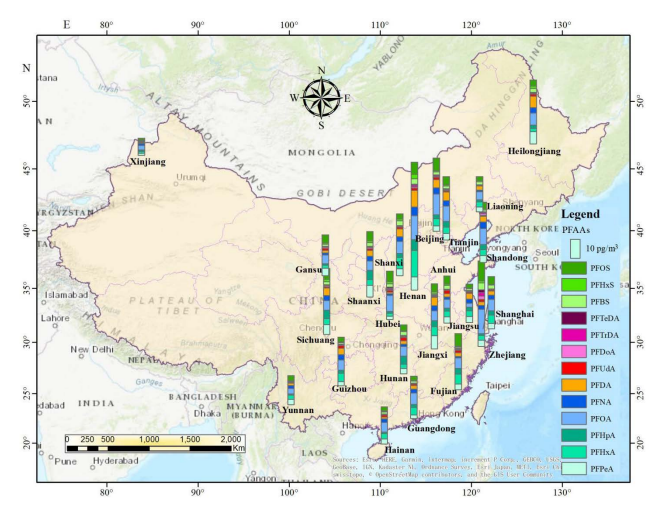


Fig. 2. The spatial distributions of PFAAs in China (annual average of PFAAs, created by ArcGIS 10.4).

3.4 Geographical distributions transport pathway

The PFAAs variations in the atmosphere depended on their local source emissions as well as regional atmosphere transportation. In order to give readers a direct impression of factors affecting the geographical variations of PFAAs in China, here we analyzed PFAAs variations along three pathway transects and one coastal line to determine how PFAAs distribute spatially.

As shown in Figure 3a, PFAAs concentrations were enriched in southeastern areas (40.58–47.17 pg/m³) at low altitudes (2–30 m), but relatively low abundances (12.31–29.44 pg/m³) existed in the northwestern part of China (397–1,517 m in altitude). As discussed above, the EC areas (e.g. Fujian) were the most intensively industrialized regions, direct emissions from PFAAs manufacturing processes would enhance their atmospheric abundances. However, high altitudes existed in NW areas would have a blocking effect to the transportation of PFAAs from eastern polluted areas.

In terms of the SW-NE transect (Figure 3b), Yunnan and Liaoning showed much lower PFAAs concentrations (20.88 and

24.99 pg/m³) than other areas (44.76–52.58 pg/m³). Notably, a steady increasing trend of PFAAs concentrations existed across the W–E transect (Figure 3c), which escalated from 20.88 pg/m³ in Yunnan to 61.68 pg/m³ in Zhejiang. The composition profiles of PFAAs along this transect differed from each other; for instance, PFOA occupied 28.5% of the total PFAAs in Zhejiang, while it only accounted for 15.6%–21.8% in other areas. Note that PFAAs released from point sources would be eliminated by deposition, degradation, or dilution during transportation in the atmosphere, e.g., PFOA could decrease by ~90% within 5 km of its point source (Chen et al., 2018). However, the long range transport of PFAAs bounded with particles also have been explored in previous research (Pickard et al., 2018). As illustrated in Figure S5, the 48 hours back trajectories were generally associated with air masses originating from the surrounding areas of the sampling locations, the trajectories which overlapped with urban areas in Zhejiang, Jiangxi and Shanghai, which confirmed that the air mass origins was a driving factor for PFAAs variation.

Interestingly, with the exclusion of the site directly affected by surrounding sources in Zhejiang, PFAAs were rather

uniformly distributed among the coastal areas, with concentrations ranging from 24.92–45.76 pg/m³ (Figure 3d). Excluded industrial and domestic emissions as well as secondary formation, the PFAAs containing sea spray aerosols could contribute the variations of PFAAs in coastal atmosphere (Cai et al., 2012b; Pickard et al., 2018).

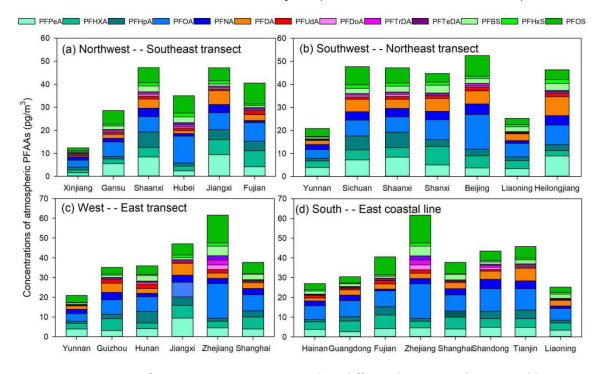


Fig. 3. Transects of PFAAs concentrations across three different directions and one coastal line

3.5 Source identification

Understanding the sources of PFAAs and their corresponding importance would enable elucidation of the levels of

PFAAs in the environment. As discussed above, the observations from tempo-spatial variations of PFAAs suggest that several factors may have a combined effect on the variations of PFAAs. Hence, a PMF model was adopted to extract the potential factors affecting PFAAs variations, and four sources were extracted in this study (see Figure 4). High percentages (~90.0%) of PFPeA and PFBS were found in factor 1, and were moderately loaded with PFOS (62.6%). Three major types of PFOS-related chemicals; namely PFOS salts, PFOS substances and PFOS polymers, are used in industrial products in China (Xie et al., 2013). PFOS salts are usually used in metal plating, firefighting foams, and pesticides, while PFOS substances are adopted in paper treatment and the semiconductor industry. PFOS polymers are employed for textile and leather treatment. These PFOS-related products would lead to direct emissions of PFOS during their industrial and domestic activities. PFPeA and PFBS are the main substitutes for long-chain PFAAs in China, which would release as impurities or by-products when manufacturing PFOS-based products (Liu et al., 2017). Hence, this factor was regarded as the direct source of PFOS-based products. This was consistent with the spatial observations that high PFOS concentrations were shown in Zhejiang, Fujian, Guangdong, and Shanghai, where manufacturing facilities are distributed. Factor 2 was characterized by PFHxA, PFOA, PFNA, and PFDA, each representing over 60.0% of their explained variations. Their rather strong positive correlations (r=0.54–0.84, p<0.01) suggested that they may have originated from a similar source (Table S6). PFOA was considered as the marker for the emulsification of plastics, rubber products, flame retardants for textiles, paper surface treatments, fire foams and PTFE emulsifiers (Liu et al., 2015; Konstantinos et al., 2010). It has been reported that there was an increase in PFCAs emissions at the manufacturing sites of PFOA-based products in China between 2002 and 2012 due to a rapid increase in domestic demand and production of PFOA-related products (Wang et al., 2014). PFNA and its derivatives have similar physicochemical properties to PFOA and its derivatives, and both can be emitted through exhaust gases. The PFNA-based production was found to be related to polyvinylidene fluoride (PVDF) production, and it has been suggested that PVDF production increased in China after 2008 (Wang et al., 2014). Therefore, factor 2 represents direct sources of PFOA-based and PFNA-based products. The compositions of factor 3 were characterized by a high loading of PFHpA and PFHxS, with loading factor values of 84.9% and 81.7%, respectively. The historical production and uses of PFHpA and its derivatives remain unidentified. Factor with PFHxS alone did not indicate a specific source, so this factor may be classified as an unknown source, which may be affected by atmosphere air mass transport, sea aerosol bursting and/or other origins. The final factor was dominated by PFUdA, PFDoA, PFTrDA, and PFTeDA, with loading factor values larger than 80%. These long-chain PFAAs (C11-C14) analogues have been interpreted as degradation products of fluorotelomer-based products in previous research (Liu et al., 2017; Wang et al., 2014; Thackray and Selin, 2017). Based on the life-cycle

311

312

313

314

315

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

usage and release from fluorotelomer and other fluorinated products, the global cumulative estimation of PFUdA, PFDoA, PFTrDA, and PFTeDA from quantified sources was estimated to be 9–230 tons in the period of 2003–2015, and projected to be between 0–84 tons between 2016–2030 (Wang et al., 2014). It was reported that the manufacturing of fluorotelomer–based substances would increase in China. In addition, these four analogues showed apparent positive correlations to each other (r =0.59–0.79, p<0.01). Thus, this factor was explained as the degradation products of fluorotelomer–based products, which could be proven by their higher abundances caused by an enhanced atmospheric oxidation ability in the summer than other seasons.

Direct emission sources, including PFOS–based products, PFOA–based products, and PFNA–based products were

Direct emission sources, including PFOS-based products, PFOA-based products, and PFNA-based products were estimated to represent 62.7% of the total PFAAs sources. Indirect sources of degradation products of fluorotelomer-based products played a minor role, contributing 15.5%, and there are 21.8% of variances that could still not be explained and need further detailed investigation. This source apportionment result was similar to one recent piece of research that found that industrial PFOA emissions were the major sources of atmospheric PFAAs in Shenzhen, China (Liu et al., 2015), and the long-distance transportation of pollutants also made a contribution.

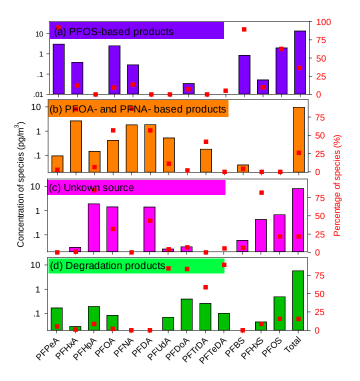


Fig. 4. Factor profiles of PFAAs extracted by the PMF model

4. Conclusion

In the present study, PFAAs were ubiquitously detected in the atmosphere across China over the length of a year. Results

indicated that the measured PFAAs in the present study were several times to several magnitudes higher than the levels conducted in most other urban locations, while far lower than the measurements implemented at point sources. In which, the C5–C14 PFCAs analogues occupied 79.6% of the total PFAAs variations, PFOA, PFHxA and PFOS ranked the top three species. Additionally, much higher abundances of PFAAs existed in winter compared with in summer. In terms of spatial distribution, the PFAAs concentrations were higher in central and eastern China, where dense residential and industrial manufacturing facilities were distributed. Correlation analysis, Hysplit backward trajectories, and PMF receptor model, have combined to suggest that the direct sources of PFOS–based, PFOA–based, and PFNA–based products made a predominant contribution to variations in PFAAs, while indirect degradation played a minor role.

Acknowledgements

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

This study was financially supported by National Key Research & Development Plan (2016YFC0200104), National Natural Science Foundation of China (No. 21577090 and No. 21777094), and China Postdoctoral Innovative Talent Support Project (BX20190169). We thank Lei Ye (Xi'an University of Architecture and Technology), Fengxia Wang (Hainan University), Linrui Jia (Beijing Normal University), Songfeng Chu (Tongji University), and other 18 volunteers, for coordinating the sampling process and for their valuable contribution to field measurement. We appreciate senior engineer Xiaofang Hu (Instrumental Analysis Center, SESE, Shanghai Jiao Tong University) for her assistance in experiment analysis.

Appendix A: Supplementary material

References

- Ahrens, L., Harner, T., Shoeib, M., Koblizkova, M., and Reiner, E. J.: Characterization of Two Passive Air Samplers for
- 377 Per- and Polyfluoroalkyl Substances, Environ. Sci. Technol., 47, 14024-14033, https://doi.org/10.1021/es4048945, 2013.
- 378 Armitage, J. M., Hayward, S. J., and Frank, W.: Modeling the Uptake of Neutral Organic Chemicals on XAD Passive Air
- 379 Samplers under Variable Temperatures, External Wind Speeds and Ambient Air Concentrations (PAS-SIM), Environ. Sci.
- 380 Technol., 47, 13546-13554, https://doi.org/10.1021/es402978a, 2013.
- Baard Ingegerdsson, F., Line Sm. Stuen, H., Raymond, O., Hanne Line, D., Merete, H., Cathrine, T., Syvert, T., Georg, B.,
- Paal, M., and Ellingsen, D. G.: Occupational exposure to airborne perfluorinated compounds during professional ski
- waxing, Environ. Sci. Technol., 44, 7723-7728, https://doi.org/10.1021/es102033k, 2010.

- Buck, R. C., Franklin, J., Berger, U., Conder, J. M., Cousins, I. T., Voogt, P. D.: Perfluoroalkyl and polyfluoroalkyl
- substances in the environment: terminology, classification, and origins. Int. Environ. Assess. Manag., 7(4), 513-541,
- 386 https://doi.org/10.1002/ieam.258, 2011.
- 387 Cai, M., Xie, Z., Moeller, A., Yin, Z., Huang, P., Cai, M., Cai M. Yang, H., Sturm, R., and He, J.: Polyfluorinated
- compounds in the atmosphere along a cruise pathway from the Japan Sea to the Arctic Ocean, Chemosphere, 87(9),
- 389 989-997 https://doi.org/ 10.1016/j.chemosphere.2011.11.010, 2012a.
- 390 Cai M., Zheo, Z., Yin Z., Ahrens L., Huang P., Cai M., Yang H., He J., Sturm R., Ebinghaus R., Xie Z.: Occurrence of
- 391 perfluoroalkyl compounds in surface waters from the North Pacific to the Arctic Ocean, Environ. Sci. Technol., 46,
- 392 661-668, https://doi.org/10.1021/es2026278, 2012b.
- Cardenas, A., Gold, D. R., Hauser, R., Kleinman, K. P., Hivert, M. F., Calafat, A. M., Ye, X., Webster, T. F., Horton, E. S.,
- and Oken, E.: Plasma Concentrations of Per- and Polyfluoroalkyl Substances at Baseline and Associations with Glycemic
- 395 Indicators and Diabetes Incidence among High-Risk Adults in the Diabetes Prevention Program Trial, Environ Health
- 396 Perspect, 125, 107001, https://doi.org/10.1289/EHP1612, 2017.
- Cassandra, R., Tom, H., K, S. J., Anita, E., Gilberto, F., Eugenia, C. L., Oscar, F., Martin, V. I., S.B., M. K., and Isabel, M.
- 398 R.: Atmospheric concentrations of new POPs and emerging chemicals of concern in the Group of Latin America and
- 399 Caribbean (GRULAC) region, Environ. Sci. Technol., 52(13), 7240-7249, https://doi.org/10.1021/acs.est.8b00995, 2018.
- 400 Chen, H., Yao, Y., Zhao, Z., Wang, Y., Wang, Q., Ren, C., Wang, B., Sun, H., Alder, A. C., and Kannan, K.: Multimedia
- 401 Distribution and Transfer of Per- and Polyfluoroalkyl Substances (PFASs) Surrounding Two Fluorochemical
- 402 Manufacturing Facilities in Fuxin, China, Environ. Sci. Technol., 52, 8263-8271, https://doi.org/10.1021/acs.est.8b00544,
- 403 2018.
- 404 De Silva, A. O.: Degradation of fluorotelomer alcohols: a likely atmospheric source of perfluorinated carboxylic acids,
- 405 Environ. Sci. Technol., 38, 3316-33121, https://doi.org/10.1021/es049860w, 2004.
- Dreyer, A., Weinberg, I., Temme, C., and Ebinghaus, R.: Polyfluorinated Compounds in the Atmosphere of the Atlantic
- 407 and Southern Oceans: Evidence for a Global Distribution, Environ. Sci. Technol., 43, 6507-6514, https://doi.org/
- 408 10.1021/es9010465, 2009.
- 409 Fang, X., Wang, Q., Zhao, Z., Tang, J., Tian, C., Yao, Y., Yu, J., and Sun, H.: Distribution and dry deposition of
- 410 alternative and legacy perfluoroalkyl and polyfluoroalkyl substances in the air above the Bohai and Yellow Seas, China,
- 411 <u>Atmos. Environ., 192, 128-135, https://doi.org/10.1016/j.atmosenv.2018.08.052, 2018.</u>
- Gewurtz, S. B., Backus, S. M., Silva, A. O., De, Lutz, A., Alain, A., Marlene, E., Susan, F., Melissa, G., Paula, G., and
- 413 Tom, H.: Perfluoroalkyl acids in the Canadian environment: multi-media assessment of current status and trends, Environ.

- 414 Int., 59, 183-200, https://doi.org/10.1016/j.envint.2013.05.008, 2013.
- 415 Han, D., Fu, Q., Gao, S., Li, L., Ma, Y., Qiao, L., Xu, H., Liang, S., Cheng, P., Chen, X., Zhou, Y., Yu, J. Z., and Cheng,
- 416 J.: Non-polar organic compounds in autumn and winter aerosols in a typical city of eastern China: size distribution and
- 417 impact of gas-particle partitioning on PM2.5 source apportionment, Atmos. Chem. Phys., 18, 9375-9391, https://doi.org/
- 418 10.5194/acp-18-9375-2018, 2018.
- 419 Han, D., Fu, Q., Gao, S., Zhang, X., Feng, J., Chen, X., Huang, X., Liao, H., Cheng, J., and Wang, W.: Investigate the
- 420 impact of local iron-steel industrial emission on atmospheric mercury concentration in Yangtze River Delta, China,
- 421 Environ. Sci. Pollut. Res., 26(6), 5862–5872, https://doi.org/10.1007/s11356-018-3978-7, 2019.
- 422 Hu, X. C., Andrews, D. Q., and Lindstrom, A. B.: Detection of Poly- and Perfluoroalkyl Substances (PFASs)in U.S.
- 423 Drinking Water Linked to Industrial Sites, Military Fire TrainingAreas, and Wastewater Treatment Plants, Environ. Sci.
- 424 Technol. Lett., 3, 344-350, https://doi.org/10.1021/acs.estlett.6b00260, 2016.
- Jian, J. M., Guo, Y., Zeng, L., Liu, L. Y., Lu, X., Wang, F., and Zeng, E. Y.: Global distribution of perfluorochemicals
- 426 (PFCs) in potential human exposure source—A review, Environ. Int., 108, 51-62, https://doi.org/, 2017.
- 427 Johansson, N., Fredriksson, A., and Eriksson, P.: Neonatal exposure to perfluorooctane sulfonate (PFOS) and
- 428 perfluorooctanoic acid (PFOA) causes neurobehavioural defects in adult mice, Neurotoxicology, 29, 160-169,
- 429 https://doi.org/10.1016/j.neuro.2007.10.008, 2008.
- 430 Karásková, P., Codling, G., Melymuk, L., and Klánová, J.: A critical assessment of passive air samplers for per- and
- 431 polyfluoroalkyl substances, Atmos. Environ., 185, 186-195, https://doi.org/10.1016/j.atmosenv.2018.05.030, 2018.
- 432 Konstantinos, P., Cousins, I. T., Buck, R. C., and Korzeniowski, S. H.: Sources, fate and transport of
- 433 perfluorocarboxylates, Environ. Sci. Technol., 40(1), 32-44, https://doi.org/10.1002/chin.200611255, 2010.
- 434 Krogseth, I. S., Xianming, Z., Ying, D., Lei, Frank, W., and Knut, B.: Calibration and application of a passive air sampler
- 435 (XAD-PAS) for volatile methyl siloxanes, Environ. Sci. Technol., 47, 4463-4470, https://doi.org/10.1021/es400427h,
- 436 2013.
- 437 Lai, F. Y., Rauert, C., Gobelius, L., and Ahrens, L.: A critical review on passive sampling in air and water for per- and
- 438 polyfluoroalkyl substances (PFASs), TrAC Trends Anal. Chem., Available online 23 Nov. 2018, https://doi.org/
- 439 10.1016/j.trac.2018.11.009, 2018.
- Li, J., Vento, S. D., Schuster, J., Gan, Z., Chakraborty, P., Kobara, Y., and Jones, K. C.: Perfluorinated Compounds in the
- 441 Asian Atmosphere, Environ. Sci. Technol., 45, 7241-7428, https://doi.org/10.1021/es201739t, 2011.
- Lindstrom, A. B., Strynar, M. J., and Libelo, E. L.: Polyfluorinated Compounds: Past, Present, and Future, Environ. Sci.
- 443 Technol., 45, 7954-7961, https://doi.org/10.1021/es2011622, 2011.

- 444 Liu, B., Zhang, H., Yao, D., Li, J., Xie, L., Wang, X., Wang, Y., Liu, G., and Yang, B.: Perfluorinated compounds (PFCs)
- in the atmosphere of Shenzhen, China: Spatial distribution, sources and health risk assessment, Chemosphere, 138,
- 446 511-518, https://doi.org/10.1016/j.chemosphere.2015.07.012, 2015.
- 447 Liu, Z., Lu, Y., Wang, P., Wang, T., Liu, S., Johnson, A. C., Sweetman, A. J., and Baninla, Y.: Pollution pathways and
- 448 release estimation of perfluorooctane sulfonate (PFOS) and perfluorooctanoic acid (PFOA) in central and eastern China,
- 449 Sci. Total Environ., 580, 1247-1256, https://doi.org/10.1016/j.scitotenv.2016.12.085, 2017.
- 450 Martin, J. W., Ellis, D. A., Mabury, S. A., Hurley, M. D., and Wallington, T. J.: Atmospheric chemistry of
- 451 perfluoroalkanesulfonamides: kinetic and product studies of the OH radical and Cl atom initiated oxidation of N-ethyl
- 452 perfluorobutanesulfonamide, Environ. Sci. Technol., 40, 864-872, https://doi.org/10.1021/es051362f, 2006.
- 453 Melymuk, L., Bohlin, P., Sáňka, O., Pozo, K., and Klánová, J.: Current challenges in air sampling of semivolatile organic
- 454 contaminants: sampling artifacts and their influence on data comparability. Environ. Sci. Technol., 48(24), 14077-91,
- 455 https://doi.org/10.1021/es502164r, 2014.
- Okeme, J. O., Saini, A., Yang, C., Zhu, J., Smedes, F., and Klànovà, J.: Calibration of polydimethylsiloxane and
- 457 XAD-pocket passive air samplers (PAS) for measuring gas- and particle-phase SVOCs. Atmos. Environ., 143, 202-208,
- 458 https://doi.org/10.1016/j.atmosenv.2016.08.023, 2016.
- 459 Pickard H. M., Criscitiello A. S., Spencer C., Sharp M. J., Muir D.C. G., De Silva A. O., Young C. J.: Continuous
- 460 non-marine inputs of per- and polyfluoroalkyl substances to the High Arctic: a multi-decadal temporal record, Atmos.
- 461 Chem. Phys., 18, 5045–5058, https://doi.org/10.5194/acp-18-5045-2018, 2018.
- 462 Rauert, C., Harner, T., Schuster, J. K., Eng, A., Fillmann, G., Castillo, L. E., Fentanes, O., Villa, M. I., Miglioranza, K.,
- and Moreno, I. R.: Atmospheric Concentrations of New Persistent Organic Pollutants and Emerging Chemicals of
- 464 Concern in the Group of Latin America and Caribbean (GRULAC) Region, Environ. Sci. Technol., 52(13), 7240-7249,
- 465 https://doi.org/ 10.1021/acs.est.8b00995, 2018.
- 466 Sedlak, M. D., Benskin, J. P., Wong, A., Grace, R., and Greig, D. J.: Per- and polyfluoroalkyl substances (PFASs) in San
- 467 Francisco Bay wildlife: Temporal trends, exposure pathways, and notable presence of precursor compounds,
- 468 Chemosphere, 185, 1217-1226, https://doi.org/10.1016/j.chemosphere.2017.04.096, 2017.
- 469 Thackray, C. P., Selin N. E.: Uncertainty and variability in atmospheric formation of PFCAs from fluorotelomer
- 470 precursors, Atmos. Chem. Phys., 17, 4585–4597, https://doi:10.5194/acp-17-4585-2017, 2017.
- Tian, Y., Yao, Y., Chang, S., Zhao, Z., Zhao, Y., Yuan, X., Wu, F., and Sun, H.: Occurrence and Phase Distribution of
- Neutral and Ionizable Per- and Polyfluoroalkyl Substances (PFASs) in the Atmosphere and Plant Leaves around Landfills:
- 473 A Case Study in Tianjin, China, Environ. Sci. Technol., 52, 1301-1310, https://doi.org/10.1021/acs.est.7b05385, 2018.

- Tian, Y., Zhou, Y., Miao, M., Wang, Z., Yuan, W., Liu, X., Wang, X., Wang, Z., Wen, S., and Liang, H.: Determinants of
- 475 plasma concentrations of perfluoroalkyl and polyfluoroalkyl substances in pregnant women from a birth cohort in
- 476 Shanghai, China, Environment International, 119, 165-173, https://doi.org/10.1016/j.envint.2018.06.015, 2018.
- Wang, Q. W., Yang, G. P., Zhang, Z. M., and Jian, S.: Perfluoroalkyl acids in surface sediments of the East China Sea,
- 478 Environ. Pollut., 231, 59-67, https://doi.org/10.1016/j.envpol.2017.07.078, 2017.
- Wang, Z., Cousins, I. T., Scheringer, M., Buck, R. C., and Hungerbühler, K.: Global emission inventories for C4-C14
- 480 perfluoroalkyl carboxylic acid (PFCA) homologues from 1951 to 2030, Part I: production and emissions from
- 481 quantifiable sources, Environ. Int., 70, 62–75, https://doi.org/10.1016/j.envint.2014.04.013, 2014.
- 482 Wang X., Schuster J., Jones K. C., Gong P.:Occurrence and spatial distribution of neutral perfluoroalkyl substances and
- 483 cyclic volatile methylsiloxanes in the atmosphere of the Tibetan Plateau, Atmos. Chem. Phys., 18, 8745–8755,
- 484 https://doi.org/10.5194/acp-18-8745-2018, 2018.
- Wong, F., Shoeib, M., Katsoyiannis, A., Eckhardt, S., Stohl, A., Bohlinnizzetto, P., Li, H., Fellin, P., Su, Y., and Hung, H.:
- 486 Assessing temporal trends and source regions of per- and polyfluoroalkyl substances (PFASs) in air under the Arctic
- 487 Monitoring and Assessment Programme (AMAP), Atmos. Environ., 172, 65-73, https://doi.org/
- 488 10.1016/j.atmosenv.2017.10.028, 2018.
- 489 Xie, S., Wang, T., Liu, S., Jones, K. C., Sweetman, A. J., and Lu, Y.: Industrial source identification and emission
- 490 estimation of perfluorooctane sulfonate in China, Environ. Int., 52, 1-8, https://doi.org/10.1016/j.envint.2012.11.004,
- 491 2013.
- 492 Yao, Y., Chang, S., Sun, H., Gan, Z., Hu, H., Zhao, Y., and Zhang, Y.: Neutral and ionic per- and polyfluoroalkyl
- substances (PFASs) in atmospheric and dry deposition samples over a source region (Tianjin, China), Environ Pollut, 212,
- 494 449-456, https://doi.org/10.1016/j.envpol.2016.02.023, 2016a.
- 495 Yao, Y., Sun, H., Gan, Z., Hu, H., Zhao, Y., Chang, S., and Zhou, Q.X.: A Nationwide Distribution of Per- and
- 496 Polyfluoroalkyl Substances (PFASs) in Outdoor Dust in Mainland China From Eastern to Western Areas, Environ. Sci.
- 497 Technol., 50(7), 3676-3685, https://doi.org/10.1021/acs.est.6b00649, 2016b.
- 498 Zhen, W., Xie, Z., Möller, A., Mi, W., Wolschke, H., and Ebinghaus, R.: Atmospheric concentrations and gas/particle
- 499 partitioning of neutral poly- and perfluoroalkyl substances in northern german coast. Atmos. Environ., 95(1), 207-213,
- 500 https://doi.org/10.1016/j.atmosenv.2014.06.036, 2014.