1 Odds and ends of atmospheric mercury in Europe and over northern Atlantic Ocean: Tem-2 poral trends of 25 years of measurements.

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- 17 Manuscript aim:

18 To determine the atmospheric mercury trend on a continental scale and evaluate the driving factor of 19 the downward trend in mercury in the Northern Atlantic and Europe. Also, to assess the time

- 20 variability in the light of atmospheric transport patterns, and regional sources.
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23 Abstract

24 The Global Monitoring Plan of the Minamata Convention on Mercury was established to generate 25 long-term data necessary for evaluating the effectiveness of regulatory measures at a global scale. 26 After 25 years monitoring (since 1995), Mace Head is one of the atmospheric monitoring stations with 27 the longest mercury record, and has produced sufficient data for the analysis of temporal trends of 28 Total Gaseous Mercury (TGM) in Europe and the Northern Atlantic. Using concentration-weighted 29 trajectories for atmospheric mercury measured at Mace Head as well as other five locations in Europe, 30 Amderma, Andøya, Villum, Waldhof and Zeppelin we identify the regional probabilistic source 31 contribution factor and its changes for the period of 1996 to 2019.

32 Temporal trends indicate that concentrations of mercury in the atmosphere in Europe and the 33 Northern Atlantic have declined significantly over the past 25 years, at a non-monotonic rate 34 averaging of 0.03 ng m⁻³ year⁻¹. Concentrations of TGM at remote marine sites were shown to be 35 affected by continental long-range transport, and evaluation of reanalysis back-trajectories display a 36 significant decrease of TGM in continental air masses from Europe in the last two decades. In addition, 37 using the relationship between mercury and other atmospheric trace gases that could serve as a 38 source signature, we perform factorization regression analysis, based on positive rotatable 39 factorization to solve probabilistic mass function. We reconstructed atmospheric mercury 40 concentration and assessed the contribution of the major natural and anthropogenic sources. The 41 results reveals that the observed downward trend in the atmospheric mercury is mainly associated42 with a factor with a high load of long-lived anthropogenic species.

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45 **1** Introduction

46 Mercury is a toxic pollutant of crucial concern to public health globally. Due to its neurotoxicity, 47 bioaccumulation, and long-range atmospheric transport, mercury was added to the priority list of 48 several international agreements and conventions dealing with environmental protection, including 49 the Minamata Convention on Mercury (e.g. Driscoll et al., 2013). Following the entry-into-force of the 50 Stockholm Convention (SC) in 2004 accompanied by the Minamata convention in 2013 to restrict 51 releases of mercury and its compounds to the environment, a Global Monitoring Plan was devised to 52 evaluate the effectiveness of regulatory measures at regional and global scales. At this time, regions 53 such as Western Europe and North America have already established monitoring networks for 54 mercury in air and precipitation some of which have been in operation since the 1990s (Schmeltz et 55 al., 2011; Gay et al., 2013; EMEP, 2020; www.gmos.eu; www.gos4m.org).

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57 During the past decades, atmospheric mercury concentrations in the Northern Hemisphere decreased 58 substantially (Slemr et al., 2003; Cole et al., 2014; Steffen et al., 2015; Weigelt et al., 2015; Weiss-59 Penzias et al. 2016; Marumoto et al., 2019; Custodio et al. 2020). This downward trend has been 60 attributed to decreasing emissions from the North Atlantic Ocean due to decreasing mercury 61 concentrations in subsurface water (Soerensen et al., 2012) and more recently to decreasing global 62 anthropogenic emissions mainly due to the decline of mercury release from commercial products 63 (Horowitz et al., 2014) and the changes of Hg^0/Hg^{2+} speciation in flue gas of coal-fired utilities after 64 implementation of NOx and SO₂ emission controls (Zhang et al., 2016). Mercury uptake by terrestrial 65 vegetation has also been recently proposed as a contributor to the downward trend (Jiskra et al., 66 2018).

As reported by Lyman et al. (2020), the mercury emission to the atmosphere is continuously changing.
Its monitoring is needed to track the trends, identify persistent and new sources, and assess the
efficacy of mercury pollution control policies.

In a 5-year source apportionment study, Custodio et al. (2020) show that a factor with high load of long lived anthropogenic atmospheric species could explain the decrease of TGM at Mace Head. This decrease is consistent with a decrease in the anthropogenic mercury emissions inventory in Europe and North America (Horowitz et al. 2014). Wu et al., (2016) estimated that China's emissions also decreased since 2012 which could have a hemispheric effect. However, the downward trend of global 75 anthropogenic mercury emissions needs to be confirmed by atmospheric observations, and a long-76 term evaluation of the time series of still not unknown sources and its implication should be assessed. 77 This study reports continuous long-term temporal trends of TGM in the Northern Atlantic, Arctic, and 78 Europe, reporting mercury atmospheric concentrations at Mace Head (1995-2019), Amderma 2001-79 2017), Andøya (2010-2019), Villum (1999-2019), Waldhof (2005-2019), and Zeppelin (2000-2019). 80 Here, we combine a long-time series of atmospheric mercury observed at these sites with calculated 81 120-hour reanalysis backward trajectories in order to investigate transport and long-term changes in 82 concentration patterns on the regional scale. 83 This paper aims to evaluate the TGM trend on a continental scale and the contribution of the baseline 84 factor as a driver of the downward trend in mercury for the Northern Atlantic and Europe.

Based on long-range Lagrangian reanalysis backward trajectories and receptor-modelling, we investigate the trends and sources of mercury in the atmosphere, assessing the inter-annual variability on the light of atmospheric transport patterns and changes in the regional emissions. In addition, we exploit atmospheric mercury temporal variability, which can be used as additional constraints to improve the ability of models to predict the cycling of mercury in the atmosphere.

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91 2 Experimental

Sampling sites

95 Data from six sites in Europe and Greenland with the longest records of atmospheric mercury 96 concentrations were selected for this study: Mace Head (data available 1995 - 2019), Zeppelin (2000 97 - 2019), Waldhof (2006 - 2019), Villum (2008 - 2019), Andøya (2010 - 2019), and Amderma (2001 -98 2013). Mace Head and Waldhof are mid-latitude stations, Zeppelin, Amderma, and Villum can be 99 classified as Arctic ones. Andøya, though at latitude comparable to that of Amderma, is behaves more 100 like a mid-latitude station because the ocean around it is ice free for most of the year. At all sites 101 mercury had been measured by a Tekran instrument (Tekran Inc, Toronto, Canada), more details will 102 be given at the end of the section

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The Mace Head Global Atmosphere Watch (GAW) Station (53°20' N and 9°32W, 8 m above sea level; air-sampling inlet 18 m a.s.l.) is located on the west coast of Ireland on the shore of the North Atlantic Ocean, offering ideal conditions to evaluate both natural and anthropogenic pollutants in oceanic and continental air masses as described by Stanley et al. (2018). The station was part of the GMOS network and mercury measurements are described in detail by Weigelt et al. (2015).

- 110The Zeppelin GAW station is located on the ridge of the Zeppelin Mountain (78°54`N, 11°52`E) at 474111m a.s.l., about 2 km from Ny Ålesund on the west coast of Spitsbergen which is the largest of the
- 112 Svalbard Islands. Mercury measurements are described by Aspmo et al. (2005).
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Waldhof (52°48'N, 10°45'E, 74 m a.s.l.) is a rural background site located in the northern German
lowlands in a flat terrain, 100 km south-east of Hamburg., The site and analytical method are described
in detail by Weigelt et al. (2013).

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Villum Research Station is located at the military outpost Station Nord. It is located in the furthermost northeastern corner of Greenland on the north–south oriented peninsula of Princess Ingeborg Halvø (81°36′ N, 16°40′ W, 25 m a.s.l.), whose northern end is a 20 × 15 km² Arctic lowland plain. The Air Observatory is located 2 km south of the central complex of Station Nord that is manned year-round by 5 soldiers. The monitoring site is upwind of the dominant wind direction for Station Nord and thus any effect of local pollution is minimized. Atmospheric measurements at Villum are described in detail by Skov et al. (2004 and 2020).

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Andøya observatory (69.3°N, 16°E, 380 m a.s.l.) is situated a few hundred meters away from ALOMAR (Arctic Lidar Observatory for Middle Atmosphere Research), which is located at the west coast on a mountain at the island Andøya in Northern Norway. ALOMAR is part of Andøya Space Center. More details about measurements at Andøya are available in Berg et al. (2008).

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Amderma Polar Station is located near the Amderma settlement of the Arkhangelsk Arctic region of Russia near the coast of the Kara Sea (69°43`N, 61°37`E, 49 m a.s.l.; Yugor Peninsula, Russia). Gaseous mercury has been measured since 2001 until 2017. The site and the mercury measurements are described by Pankratov et al. (2013).

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At all sites mercury was measured using Tekran 2537 A and/or B instrument (Tekran Inc, Toronto, Canada, mostly Model A, at Mace Head and Villum also Model B), an automated dual-channel, single amalgamation, cold vapor atomic fluorescence (CVAFS) analyzer. The instrument has two gold cartridges. While mercury is collected on one of them during the sampling period, the other is being analyzed by thermodesorption and CVAFS detection. The functions of the cartridges are then alternated, allowing for quasi-continuous measurement. The instruments are usually protected by an upstream PTFE filter against dust and aerosols.

143 As discussed by Slemr et al. (2016), gaseous oxidized mercury (GOM) compounds are collected on 144 the gold cartridges and were found to be converted to elemental mercury (GEM) probably during 145 the thermodesorption. The instrument is thus able to measure total gaseous mercury (TGM) pro-146 vided that GOM compounds reach the cartridges. This is frequently not the case because the GOM 147 compounds are sticky and can thus be removed on the way from the inlet to the cartridges (Lyman 148 et al., 2020). The instruments are usually protected by an upstream PTFE filter (mostly 0.2 μm, 0.4 149 μm at Zeppelin, 0.45 μm at Andøya) against dust and aerosols. Additional soda-lime filters are fre-150 quently used to remove free halogens that can shorten the lifetime of the gold cartridges (GMOS 151 Standard Operating Procedure, 2019) and were implemented at Villum, Amderma, Zeppelin, and 152 Andøya. They are suspected to capture GOM although this has not been adequately tested so far 153 (Gustin et al., 2021). Sea salt on the walls of the sampling tubing and on the PTFE filter at coastal 154 stations, such as Mace Head, Andøya, Amderma, and possibly Zeppelin, is also likely to remove 155 GOM. We conclude that GEM is being measured at Mace Head (Weigelt et al., 2015), Villum (Skov 156 et al., 2020), Andøya, Amderma, and Zeppelin (Durnford et al., 2010), Waldhof (Weigelt et al., 157 2013). We thus treat all data as GEM. All instruments have been operated according to the stand-158 ard operating procedures (Steffen and Schroeder, 1999; GMOS Standard Operating Procedure, 159 2019). The instruments at Villum, Zeppelin, and Andøya were run with 5 min resolution at a sam-160 pling flow rate of 1.5 L min⁻¹. At Waldhof and Mace Head the temporal resolution was 15 min and 161 at Amderma 30 min. The detection limit of the Tekran

Speciated mercury measurements made at Waldhof between 2009 and 2011 provided median concentrations of 6.3 pg m⁻³ for PBM and 1.0 pg m⁻³ for GOM while the median GEM concentration was 1.6 ng m⁻³, representing >99,5% of the TGM (Weigelt et al., 2013). GOM measurements using Tekran speciation system are considered to be underestimated (Jaffe et al., 2014; Lyman et al. 2020). Other speciation measurements show that with the exception of polar depletion events and upper troposphere, GEM is the dominant form of atmospheric mercury, accounting mostly for more than 95% of the TGM (Mao et al., 2016)

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• Back-Trajectory Analysis, Concentration-weighted trajectories, and probability mass function models.

173 To evaluate the spatial coverage and sources of air sampled at the six stations, three dimensional 174 reanalysis air mass back-trajectories at an arrival height of 50 m and 500 m above ground were 175 calculated site 120 h HYSPLIT (v.4.2.0, at each for using 176 NOAA<u>https://www.arl.noaa.gov/hysplit/hysplit/</u>) as described by Stein et al. (2015). Two trajectories 177 were calculated per day, each representing an average trajectory for the period of 12 h. All individual back-trajectories generated by HYSPLIT were converted to text shape files and imported into R (R
 Project for Statistical Computing), merged with concentration files and used for spatial analysis. To
 account for the speed and atmospheric residence time of air masses, each continuous back-trajectory
 line was transformed into 120 hourly points.

182 Concentration-weighted trajectories (CWT), is an approach which can be used to indicate the proba-183 bility of a grid cells contribution to pollution events (Cheng at al. 2013). It is based on a statistical 184 model and can incorporate meteorological information in its analysis scheme to identify the average 185 concentration in areas for pollutants based on a conditional probability that an air parcel that passed 186 through a cell with a gradient concentration displays a high concentration at the trajectory endpoint 187 (Ashbaugh et al. 1985, Byčenkienė, et al. 2014). The CWT obtained at this study are a function of 188 average mercury concentrations that were obtained every 12 h and of the residence time of a trajec-189 tory in each grid cell. The 12-hour trajectory segment endpoints for each back trajectory that corre-190 sponds to each 12 h TGM, or GEM, were retained. For a 120-hour trajectory duration, 84 trajectory 191 segment end points were calculated. This transformation of trajectories into hourly segments allowed 192 the subsequent application of a kernel density tool to the combined back-trajectory air mass points 193 from all sampling sites in order to create a density map of the continental concentration and spatial 194 coverage of concentration airflows sampled at the sampling site over the course of an entire year. 195 Seasonal back-trajectory maps were also generated for evaluation of potential seasonal changes in 196 the coverage and sources of airflows (with seasons defined as summer (June, July, and August), au-197 tumn (September, October, and November), winter (December, January, and February), and spring 198 (March, April, and May).

199 The source apportionment for Mace Head was performed based on the mass conservation principle 200 with the inclusion of potential rotated infinity matrices transformation producing factors with 201 chemical profile signed by tracer species linked to its source. The full description of PMF and its 202 reconstruction consideration, chemical species considered, uncertainties, and constraining of factors 203 are presented in Custodio et al. (2020). In this study, the PMF was applied to the Mace Head daily 204 data. The species considered in the factorization and their mass loaded in each factor are displayed in 205 Figure 3S in the article supplement section. In addition, the reconstructed gaseous mercury and the 206 observation displayed an r^2 of 0.9949. The mercury mass solved by factorization agree into the 207 10/90th percentile quantile regression, as shown in Figure 4S in the supplement of the article.

In this study, the assessment was performed on annual bases, the concentrations in grid cells were calculated by counting the average concentration of trajectory segment end points that terminate within each cell as described by Byčenkienė, et al. (2014) and Tang et al. (2018).

- 212 **3** Results and discussion
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In this section, we present the time series and trends of GEM concentrations from a data set covering the period from February 1996 to December 2019 (Mace Head), July 2001 to March 2017 (Amderma), February 2000 to December 2019 (Zeppelin), January 2006 to December 2019 (Waldhof), from January 2004 to December 2019 (Andøya), and from June 1999 to December 2019 (<u>Vil-</u> <u>lum). At Villum the measurements covered only 6 months (spring, summer, and early autumn) in</u> <u>1999 – 2002, and no measurements are available for the years 2003 – 2008 (Skov et al., 2020).</u> The data are summarized in in Figure 1.

221 GEM concentrations and their frequency distributions shown in Figure 1 display distinct differences 222 between the stations. GEM concentrations at Villum, Amderma, and Zeppelin decrease frequently 223 to values near zero (minima of 0.0, 0.0, and 0.1 ng m⁻³ at Villum, Amderma, and Zeppelin, respec-224 tively) and their frequency distribution is skewed to lower values as documented by somewhat 225 lower average than median GEM concentrations and the lowest 5th percentiles of all sites with 0.55, 0.62, and 1.04 ng m⁻³ at Villum, Amderma, and Zeppelin, respectively. The seasonal occur-226 227 rence of the polar depletion events at these three stations is characteristic for the Arctic sites with 228 ice and snow coverage (Steffen et al., 2008). The GEM frequency distribution at Zeppelin is less 229 skewed than at Villum and Amderma perhaps because of the Zeppelin altitude of almost 500 m asl, 230 which is above the layer with most intensive halogen chemistry within the first 100 – 200 m above 231 snow (Tackett et al., 2007).

The distribution of GEM concentrations at Waldhof, a mid-latitude station in Central Europe, is on the contrary skewed to higher values because of frequent events with local and regional pollution (Weigelt et al., 2013). The average and median GEM concentrations at Waldhof are the highest of all the investigated stations, and the average GEM concentration is substantially higher than median one.

237 The frequency distribution at Andøya is nearly symmetric, neither skewed to low nor to high GEM 238 concentrations although a pronounced seasonal variation can be observed. At latitude comparable to 239 that of Amderma there are no pronounced depletion events at Andøya because it is exposed to Gulf 240 stream and as such free of ice for most of the year. Events with local and regional pollution are also 241 largely missing at Andøya (95th percentile of 1.79 ng m⁻³ is lower when compared with 2.32 and 2.96 242 ng m⁻³ at Waldhof and Mace Head). GEM frequency distribution at Mace Head is similar to that at 243 Andøya and the average and median GEM concentrations are nearly the same as both stations are 244 exposed to air originating mostly from the Atlantic Ocean. Opposite to Andøya, GEM frequency 245 distribution at Mace Head is slightly skewed to higher concentration because of the local pollution and 246 occasional air transport from Europe (Weigelt et al., 2015).

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248 **3.1** Seasonal variation

249 Figure 2 shows similar seasonal variations at Mace Head, Waldhof, and Andøya with the maximum 250 GEM concentrations in late winter and early spring and the mimimum ones in late summer and early 251 autumn. Similar seasonal variation has been observed at most of the mid-latitude sites in the northern 252 hemisphere (e.g. Cole et al., 2014; Weigelt et al., 2015; Sprovieri et al, 2016, Angot et al., 2016). It is 253 usually accompanied by a summer maximum in wet deposition (Gratz et al., 2009; Prestbo and Gay, 254 2009; Zhang and Jaeglé, 2013; Sprovieri et al., 2017) which is caused by faster oxidation of Hg⁰ to Hg²⁺ 255 in summer providing more Hg²⁺ for scavenging by rain (Holmes et al., 2010; Zhang et al., 2012; Zhang 256 and Jaeglé, 2013; Horowitz et al., 2017). GEM uptake by vegetation can also contribute to summer 257 minimum of GEM concentrations at midlatitudes (Jiskra et al., 2018).

258 Seasonal variations in mercury at Amderma, Villum and Zeppelin are influenced by polar depletion 259 events in spring and the subsequent reemission of the deposited mercury from snow in summer which 260 result in pronounced GEM minima in April and May and maxima in July (Steffen et al., 2008, 2015; 261 Dommergue et al., 2010; Cole and Steffen, 2010; Cole et al. 2014; Angot et al., 2016; Skov et al. 2020). 262 A similar pattern is also observed at Alert (Cole et al. 2014). Note the larger amplitude of seasonal 263 variation at Arctic stations $(0.8 - 1.2 \text{ ng m}^{-3})$ when compared to the mid-latitude ones (0.95 - 1.07 ng)264 m^{-3}). Zeppelin has a substantially smaller amplitude of seasonal variation than Amderma and Villum, 265 probably because of its altitude as already noted in the discussion of the frequency distributions.

Andøya, although located at a comparable latitude as Amderma, is only slightly influenced by the polar
depletion events because it is ice-free for most of the year, as already mentioned.

268 Figure 2 shows density maps which are based on the seasonal mean mercury concentration associated 269 with respective trajectories which arrived synchronously at all six stations. The northern parts of the 270 spring and summer panels show over the Arctic Ocean the lowest and highest mercury concentrations, 271 respectively, which is consistent with the spring polar mercury depletion and summer emission of the 272 mercury deposited during the depletion events. The highest GEM concentrations over the middle of 273 the North Atlantic occur in winter, the lowest ones in summer and autumn which is consistent with 274 the seasonal variations at Mace Head and Andøya. High GEM levels over large part of the Europe occur 275 in all seasons. The highest concentrations by level and extension occur in winter and spring, somewhat 276 lower in summer and autumn.

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278 **3.2** Temporal trends and regional source of GEM

Figures 3 and 1S show the Kernel-regression of mercury concentrations at Mace Head, Amderma,
 Andøya, Villum, Waldhof, and Zeppelin. Both figures show a non-monotonic concentration change

282 with temporary increases to intermediate maxima at Waldhof, Zeppelin, and most pronounced at 283 Villum with a maximum in 2013. The overall trend of GEM concentrations at all sites points in 284 downward direction. Table 1 summarizes the overall trends calculated by least-square-fit (LSQF) from 285 monthly medians and compares them with those at Mace Head over the same periods of available 286 measurements. Averages of monthly medians over the same periods are also listed. Monthly medians 287 were chosen to reduce the influence of depletion events at polar stations and of pollution events at 288 midlatitude stations. Mace Head was taken as a bench mark because of the longest and most complete 289 data record. In addition, the trend at Mace Head represents the baseline trend (Weigelt et al., 2015). 290 All trends in the table are significant at >99.9% level as are the differences between the trends at the 291 sites and those at Mace Head.

292 GEM concentration at Mace Head decreased with an annual rate of -0.0244 ± 0.0011 ng m⁻³ yr⁻¹ in 25 293 years (-0.0256 \pm 0.0012 ng m⁻³ yr⁻¹ in 24 years). For different periods within these long-term 294 measurements, the decrease rate at Mace Head varied between -0.0244 and -0.0346 ng m⁻³ yr⁻¹ as 295 illustrated by Figure 3. The average GEM concentrations at Waldhof are substantially higher than 296 those at Mace Head demonstrating the continuing presence of regional emissions. The downward 297 trend at Andøya is comparable to that at Waldhof but substantially smaller than at Mace Head for the 298 period of Andøya measurements. The average GEM concentration at Andøya is somewhat higher than 299 at Mace Head.

300 Of the Arctic stations, GEM concentration at Zeppelin decreased with only -0.0087 ng m⁻³ yr⁻¹ when 301 compared to -0.279 ng m⁻³ yr⁻¹ for the same period at Mace Head. Cole et al. (2013) have reported a 302 trend of +0.002 ng m⁻³ yr⁻¹ (-0.007 to + 0.012 ng m⁻³ yr⁻¹, 95% confidence range) for Zeppelin in the 303 decade 2000 – 2009 which is consistent with the trend value presented here for 2000 – 2019. The 304 average GEM concentration of 1.57 ± 0.24 ng m⁻³ for the decade 2000 - 2009 (Cole et al., 2013) is 305 almost identical with 1.55 \pm 0.14 ng m⁻³ reported here for the years 2000 – 2019, too. A somewhat 306 higher but comparable decrease rate of -0.012 ng m⁻³ yr⁻¹ (-0.021 to 0.000 ng m⁻³ yr⁻¹, 95% confidence 307 interval) was reported for Alert for the 2000 to 2009 period (Cole et al., 2013). The average GEM 308 concentration of 1.50 ± 0.35 ng m-3 at Alert is also comparable to that of Zeppelin in the 2000 – 2009 309 period (Cole et al., 2013). Figure 3 shows at Zeppelin a broad maximum around 2006.

Based on LSQF the GEM at the Arctic stations Amderma and Villum behave differently. The downward trends of -0.0327 ± 0.0047 and -0.0409 ± 0.0072 ng m⁻³ yr⁻¹ at Amderma and Villum, respectively, are roughly comparable and both are substantially larger than those at Mace Head for the respective periods. Their trend uncertainties are substantially larger than the uncertainties at the other stations. On the other side, the average GEM concentrations at Amderma and Villum are comparable to those at Mace Head for the respective periods, albeit with substantially higher standard deviations. This is 316 partly due to the short periods with varying trend at Amderma and even a pronounced temporal 317 maximum at Villum.

318 The higher level of atmospheric mercury at Villum in 2013 is consistent with an elevated mercury level 319 over Greenland in that year, as deduced from backward trajectory analyses shown in Figure 4. Large 320 subglacial source of mercury at Greenland has been recently reported by Hawkings et al. (2021). The 321 increase of GEM at Villum in 2010 and 2013, which drives the trend up during this period, corresponds 322 to two periods of negative extreme at Arctic Oscillation (AO). The extreme on AO and North Atlantic 323 Oscillation (NAO) can enhance the mercury discharge from ice to the atmosphere. Bevis et al. (2019) 324 report an anomalous ice mass loss at Greenland in the 2010-2014 epoch. The abrupt ice melting was 325 driven mainly by changes in air temperature and solar radiation caused by atmospheric circulation 326 anomalies.

In addition, the negative phase of the summertime NAO index increases the prevalence of high pressure, clear-sky conditions, enhancing surface absorption of solar radiation and decreasing snowfall, and it causes the advection of warm air from southern latitudes into Greenland. These changes promote higher air temperatures, a more extended ablation season and enhanced melt ice (Fettweis et al. 2013). In 2014/2015, when the AO indexes again turned positive and NAO negative, significant ice loss was reestablished (Bevis et al., 2019).

333 The back trajectories of air masses calculated for each site were combined with the measured 334 concentration at a 12h time resolution. The results were used to identify possible regional sources and 335 also to assess temporal variations. Figure 4 shows that calculated air mass back-trajectories for the 336 five monitoring sites mainly reflect air masses transported from the ocean, however, they also 337 indicated elevated concentrations in continental trajectories such as from central Europe which are 338 due to anthropogenic emission sources. Despite a shift to the south that can be associated with 339 uncertainties in the Lagrangian approach, the airflow patterns and concentrations hotspot were 340 consistent with the current knowledge of geolocation of GEM sources in Europe. Figure 4 also shows 341 a high level of mercury associated with air masses coming from the northwest (Canada and Greenland) 342 during the 1997-2000 epoch, 2005, 2010, 2014 besides of 2013 already mentioned.

343 The most revealing detail in the observed trend of GEM is displayed in Figure 4, where it is noticea-

344 ble that the downward trend is ongoing on a regional scale. This decrease could represent a change

in the balance between sources and sinks of mercury in the atmosphere.

The downward trend seems to be driven by decreasing concentrations in continental Europe. This phenomenon is observed mainly after 2005 when data from Waldhof is considered. The downward trend in mercury concentration is observed in all trajectories, even in remote areas, indicated by the yellow fades to green. This phenomenon can be explained only by reductions in global atmospheric mercury sources. In addition, Figure 4 also shows that the decrease is more pronounced in the hotspot areas identified as anthropogenic sources, where the colour shifts from dark to light red in plots from 2005 to 2019.

The later downward trend at Zeppelin and Villum (Figure 3, 1S), suggests that these remote, high latitude stations are less affected by direct European continental emission.

355 The seemingly non-monotonic downward trends with inter-annual ups and downs observed in this 356 study are not well explained. However, an inspection of the Mace Head data (e.g. in Figure 3 and 4) 357 reveal that this trend is composed of two segments: one starting in 1999 and ending approximately in 358 2010 and a second one in 2014 after a biennial upward tendency. It could be premature to assume 359 that the atmospheric mercury trend can be driven simply by a political decision. However, it can be 360 seen that the two important GEM trend deflections in 1999 and 2014, coincide with COUNCIL 361 DIRECTIVE 1999/31/EC, a European Union (EU) directive that regulates waste management of landfills 362 in the EU and the mercury international treaty (Minamata Convention on Mercury) designed to 363 protect human health and the environment from anthropogenic emissions and releases of mercury 364 approved on 10 October, 2013. Continental and international environmental treaties are the result of 365 long political and societal debate and commitment to such deal could reflect an already established 366 control policy at the national level.

For example, in 1990 The United States Clean Air Act, put mercury on a list of toxic pollutants that needed to be controlled to the greatest possible extent, forcing industries that release high concentrations of mercury into the environment to install maximum achievable control technologies (MACT). In 2005, the EPA promulgated a regulation that added power plants to the list of sources that should be controlled and instituted in the nation, and in 2011 new rules for coal-fired power plants were announced by EPA (State of new Jersey, at al. 2008, *Castro Mark S., Sherwell John 2015*).

Additionally, in 2007 the European Union implemented new mercury control measures, banning
mercury in new non-electrical measuring devices, such as thermometers and barometers (Jones, H.
2007).

We note that Waldhof, a continental station close to anthropogenic sources in Europe, corroborates
 the interpretation of an anthropogenic emission driven mercury trend. This station shows a more
 pronounced GEM decrease between 2005-2010 compared to the years since then.

An accurate emissions inventory is essential for interpreting trends in atmospheric concentrations and assessing the effectiveness of mercury pollution control policies (Lyman et al., 2020). However, the trends in GEM concentrations have not always been consistent with those of global anthropogenic emissions inventories, which uncertainties ranged from -33% to 60% (Lyman et al. 2020 and references in). Besides a conundrum in the global emission inventories, which displays an upwards
 trend, as shown in Figure 5S, the Emissions Database for Global Atmospheric Research (EDGARv4.tox2,
 2018) also displays a downward trend for Europe and Northern Atlantic (Figure 05).

2hang et al. (2016) presented a revised inventory of Hg emissions for the estimation of artisanal and small-scale gold mining emissions, and, accounting for the change in Hg⁰/Hg^{II} speciation of emissions from coal-fired utilities after implementation of emission controls targeted at SO₂ and NO_x, those authors estimate a factor of 20% decrease in atmospheric emission from 1990 to 2010. As shown in Figure 05, anthropogenic emissions in Europe and Northern Atlantic decreased by 31% from 1995 to 2012 (EDGARv4.tox2, 2018); the displayed decrease was mainly associated with the industrial sectors as Chlor-alkali and combustion power.

393 Natural sources can contribute up to 40% of the atmospheric mercury budget (Pirrone, et al., 2010);
394 however, a trend on such a source is not observed or reported in the literature, so far.

395 Based on the GEM associated with each air mass trajectory, we investigated the impact of atmospheric 396 circulation on continental Europe and Northern Atlantic Ocean and observe distinct concentration 397 patterns for the ocean and continental regions. We observed for example, that air masses arriving at 398 Mace Head from central Europe show distinct trends. We compared the regional patterns of GEM 399 with other pollutants (CO, CO₂, CH₄, O₃, CHCl₃, CCl₄, and CFCs) also measured at Mace Head and find 400 that GEM shows a similar pattern concerning source location as the other species closely related to 401 anthropogenic sources. However, GEM displays a downward trend, with decreasing concentrations in 402 air masses from central Europe and England.

Figure 4 shows the concentration- weighted trajectory maps for GEM measurements over Mace Head, Amderma, Andøya, Villum, Waldhof and Zeppelin. It can be seen that the highest concentrations are almost exclusively from air masses over central Europe. Exceptions are 1997 to 2000 which indicate high levels of GEM in air masses coming from Northwest. However, it should be mentioned that CWT for this period computed only Mace Head data and Villum (1999-2000).

The results also show a lower level of GEM in air masses segments over the North Atlantic region. This
 region is constantly associated with a sink of anthropogenic pollutants.

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411 **3.3 Probability of source contribution.**

Based on our analysis so far, our hypothesis is that the mercury concentration in North Atlantic air masses is affected by the intensity of transport from important regional and global sources and also by temporal changes in these sources. For example, the high mercury concentrations observed in the late 1990's coincide with higher contributions from continental air masses. During 2001, a noticeable reduction in the Mace Head GEM concentration was observed, corresponding to a lesser influence of 417 continental European air masses. This was due not only to a lower frequency of air masses from 418 continental Europe but also lower concentration of GEM in those air masses compared to previous 419 years. A similar phenomenon was observed in the trend during 2005/2006 and 2008 to 2010 when an 420 increase and decrease of inter-annual trend corresponded to higher and lower CWT in air masses 421 coming from continental Europe (Figure 2S).

In a five-year source apportionment of mercury at Mace Head, Custodio et al. (2020) show that a factor with high load of anthropogenic species could explain downward trends of atmospheric mercury. The downward trend of that factor was associated with a reduction in emissions due to cleaner manufacturing processes involving mercury and regulations limiting the emissions from coalfired power plants since the 1980s, as well as a reduction in the release of mercury from commercial products since 1990s (Streets et al. 2011, Zhang et al., 2016).

428 Here we extend the source apportionment analysis back to 1996. The extended reconstruction of the 429 main sources of mercury back to 1996, shown in Figure 6, displays a similar apportionment pattern to 430 that reported by Custodio et al. (2020). The source apportionment indicates a baseline factor charac-431 terized by high load of anthropogenic species accounting for 65% of GEM mass. The baseline factor 432 has already been proposed as the driving factor for mercury trends at Mace Head by Custodio et al. 433 (2020). In this study, this factor displays a downward trend of 2.7 % yr⁻¹, and correlates (r =0.97) with 434 the mercury trend (Figure 7). A factor with load of anthropogenic species driving the Mace Head GEM 435 trend down by a strength of 97 % at the level of 0.001 (p-values) is also supported by Figure 4, which 436 displays a temporal decrease in mercury level in reanalysis backward trajectory.

437 One important consideration to take into account is that the baseline factor is interpreted as global 438 mercury budget from several sources which were not solved by PMF, such factor could also take into 439 account the strength of non-modulated extremes events or periodic oscillations such as ENSO as 440 speculated by Slemr et al. (2020) and references therein, those events can be a reason for increase 441 rotation in the mercury trend, imposing significance and raising the correlation.

442 The Global Mercury Assessment inventory (AMAP/UNEP, 2019) estimates a contribution of combus-443 tion sources to atmospheric mercury at 24%. In this study the combustion factor, which was indicated 444 by high load of CO, accounted for 20% of total GEM mass at Mace Head (Figure 6). A slight decreasing 445 trend was observed in this factor, which could be associated with the implementation of emission 446 controls on coal-fired utilities as proposed by Zhang at al. (2016) in a revised inventory of Hg emissions. 447 However, as reported by Custodio at al. (2020) this trend should be taken with caution since the com-448 bustion factor was fingerprinted by CO, a short-lived species (1-3 months) with significant seasonal 449 and atmospheric transport dependence. Although, the trend in the combustion factor solved by PMF

450 complies with the decrease in the emission in the sector "combustion in residential and other com-

451 bustion", provided the EDGAR inventory and showed in Figure 05.

- 452 The ocean factors account for 12% of total GEM mass at Mace Head and was identified by a high load
- 453 of CHCl₃ (Figure 5). CHCl₃ used to trace sign ocean factor, is a trace atmospheric gas originating 90%
- 454 from a natural source, being offshore seawater the largest issuer (McCulloch, 2003).
- As reported by Custodio et al. (2020) and references therein, the residence time of mercury in the ocean is substantially longer than in the atmosphere, ranging from years to decades or millennia. Human activity has substantially increased the oceanic mercury reservoir and consequently is affecting
- 458 the fluxes of mercury between the sea and atmosphere (Strode et al., 2007).
- 459 The acidification of oceans, climate change, excess nutrient inputs, and pollution are fundamentally
- 460 changing the ocean's biogeochemistry (Doney, 2010) and will certainly also influence mercury ocean-
- 461 air fluxes in a still unknown direction.

462 This study shows an upward trend in the oceanic factor after 2010, as can be seen in Figure (6), how-

- 463 ever its significance, implication and causes remain to be determined.
- 464 Due to a lack of source markers that could allow the propagation of the eigenvector from axis rotation
- to reconstruct more realistically the complexity of mercury sources, only four factors solved our fac-
- 466 torization. However, such an approach provided be a valuable method to evaluate mercury fluxes.
- 467

468 **4 Conclusion**

- A conundrum in the observed negative trend in mercury in Europe and Northern Atlantic over the past two decades is explained in this study by a decrease in anthropogenic emissions. The significant decline in concentrations of GEM over the past two decades demonstrates that regulatory measures across Europe have been successful in reducing the atmospheric concentration of this species although an extensive fossil fuel use and a legacy of stockpiles in the environment continue to pose a challenge.
- These results show the transport pattern of atmospheric mercury and reveal that a baseline factor with a high load of long-lived anthropogenic species dominates the source of mercury in the Northern Atlantic and highlight the need for continued monitoring of the GEM and its sources. This study brings a monitoring concept for mercury on a continental scale which can be extended to a Global Monitoring plan by integration of the mercury monitoring network, potentially identifying hotspot concentration areas and their change over time.
- 481 This large-scale, long-term trend data evaluation can be used for assessing the effectiveness of the482 Minamata Convention.
- 483 More specific conclusions include the following:

- 484 > Enhancement of mercury in the air masses over Greenland in summer during epochs of 485 atmospheric circulation anomalies. Mercury downward trends of $2 \pm 3\%$ yr⁻¹, $2.1 \pm 1.5\%$ yr⁻¹, $1.6 \pm 3.9\%$ yr⁻¹, $4 \pm 16\%$ yr⁻¹, $2 \pm$ 486 \triangleright 487 4 % yr⁻¹, and 3 ± 3 % yr⁻¹ at Amaderma, Andøya, Mace Head, Villum, Waldholf, and Zeppelin 488 respectively are influenced by regional sources and then biased by atmosphere transport. 489 The observed GEM downward trend at Northern Atlantic and Arctic seems to be driven by 490 decreasing in concentration in continental Europe. 491 > A baseline factor with high load of anthropogenic species drives the mercury trend down by 492 a strength of 97 % at the level of 0.001 (p-values) based on source reconstruction at Mace 493 Head. 494 Combustion sources could account for 20 % of GEM with a decreasing trend, and ocean
- 495 sources account for 12 % with a slightly increasing trend.
- 496

497 Authors Contribution:

498 DC proposed the article, processed data and wrote the article. KAP provided data and evaluated the findings.
 499 TGS provided data, support the writing and discussions. FFP provided data and participate in the discussion. IS
 500 supported the calculation in scripts, data assimilation, besides provide meteorological and Lagrangian analysis.
 501 KP supported the trajectories calculation and discussion. HS provided data and discussion in its interpretation.
 502 JB and RE endorse and supported the article preparation, respectively.

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505Table 1: Comparison of GEM trends and average concentrations at Zeppelin, Waldhof, Andøya, Amderma, and506Villum with those at Mace Head. The trends (± confidence interval at 95% level) were calculated by the least507square fit (LSQF) of monthly medians over the same months for which the measurements are available. Average508GEM concentrations were calculated as average of monthly medians over months with synchronous measure-509ments.

510

Site	Period, number of	Trend [ng m ⁻³ yr ⁻¹]		GEM average concentration	
	months			[ng m ⁻]	
		Site	Mace Head	Site	Mace Head
Mace Head	Feb 1996 – Dec 2020, 279		-0.0244 ± 0.0011		
Mace Head	Feb 1996 – Dec 2019, 267		-0.0256 ± 0.0012		
Zeppelin	Feb 2000 – Dec 2019, 222	-0.0087±0.0015	-0.0279 ± 0.0013	1.548±0.141	1.483±0.196
Waldhof	Jan 2006 – Dec 2019, 161	-0.0243±0.0025	-0.0280 ± 0.0022	1.649±0.161	1.399±0.158
Andøya	Jan 2004 – Dec 2019, 119	-0.0262±0.0023	-0.0346 ± 0.0029	1.519±0.127	1.368±0.165
Amderma	Jul 2001 – Mar 2017, 133	-0.0327±0.0047	-0.0257 ± 0.0022	1.480±0.265	1.517±0.153
Villum	Sep 2008 – Jun 2019, 111	-0.0409±0.0072	-0.0293 ± 0.0031	1.372±0.274	1.371±0.140

511





514 Figure 1: Summary of time series of GEM (ng m⁻³) measured at Mace Head, Zeppelin, Waldhof, Andøya,

- 515 Amderma and Villum on the left side. Distributions density of the measured concentrations on the left side. *The
- 516 red and blue bars on the time axis represent the missing and available data periods, respectively.
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520 Figure 2: Left panels: The density map of atmospheric mercury concentrations in different seasons. Right panels:

521 Normalized annual variation of the mercury concentrations at Arctic stations (Amderma, Villum, Zeppelin) and

522 at the mid-latitude ones (Mace Head, Waldhof, and Andøya). The shaded areas are the 95% confidence intervals

523 for the monthly mean.





Figure 3. Kernel-regression of GEM at Amderma, Andøya, Mace Head, Villum, Waldhof, and Zeppelin for the period of 2001-2013, 2010-2019, 1995-2019, 2008-2019, 2006-2019, and 2000-2019 respectively. The smooth lines and shaded areas represent the Kernel-regression at 95% significance level. The thin lines show the monthly time series of GEM after removing annual cycles with amplitudes of 0.49 ng m⁻³, 0.23 ng m⁻³, 0.17 ng m⁻³, 0.30 ng m⁻³, 22 ng m⁻³, and 0.25 ng m⁻³ respectively for Amderma, Andøya, Mace Head, Villum, Waldhof, and Zeppelin. The annual cycle was calculated based on seasonality of the time series decomposition. *An individual plot regression for each station is presented in Figure 1S.

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537 Figure 4: Concentration level (concentration-weighted trajectory) of GEM (ng m⁻³) based on the mercury

538 concentration associated to its reanalysis backward trajectory at Amderma, Andøya, Mace Head, Villum,

539 Waldhof, and Zeppelin. *The black dots show the arriving point (stations) considered for each year.





Figure 5: Time-series of Europe and North Atlantic mercury emission. Emission inventory provided by Emissions Database for Global Atmospheric Research (EDGARv4.tox2, 2018). The inventory data is available at <u>https://ed-</u> <u>gar.jrc.ec.europa.eu/dataset_tox4#sources</u>. *The time-series displays the time variability of 12 sectors reported as cement production (cement), combustion in residential and other combustion (comb), glass production (glass), artisanal and small scale gold production (gold_A), large scale gold production (gold_L), shipping emission (shipping), road transportation (tro-roa), chloralkali industry mercury cell technology (chlor), combustion in power generation and industry (ind), and solid waste incineration and agricultural waste burning (waste).



reconstruction for Mace Head from 1996 to 2019, baseline (green) combustion (red), ocean (blue) and unknown factor (grey). The smooth lines and shaded areas represent the Kernel-regression at 95% significance level. The thin lines show the monthly time series with annual cycles removed.





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