

1 Atmospheric mixing ratios of methyl ethyl ketone (2-butanone) in tropical, bo- 2 real, temperate and marine environments

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27 Abstract

28 Methyl ethyl ketone (MEK) enters the atmosphere following direct emission from vegeta-
29 tion and anthropogenic activities, as well as being produced by the gas-phase oxidation of volatile
30 organic compounds (VOCs) such as *n*-butane. This study presents the first overview of ambient
31 MEK measurements at six different locations, characteristic of forested, urban and marine envi-
32 ronments. In order to understand better the occurrence and behaviour of MEK in the atmosphere,
33 we analyse diel cycles of MEK mixing ratios, vertical profiles, ecosystem flux data, and HYSPLIT
34 back trajectories, and compare with co-measured VOCs. MEK measurements were primarily con-
35 ducted with proton transfer reaction – mass spectrometer (PTR-MS) instruments. Results from the
36 sites under biogenic influence demonstrate that vegetation is an important source of MEK. The diel
37 cycle of MEK follows that of ambient temperature and the forest structure plays an important role
38 in air mixing. At such sites, a high correlation of MEK with acetone was observed (e.g. $r^2 = 0.96$
39 for the SMEAR-Estonia site in a remote hemi-boreal forest in Tartumaa, Estonia, and $r^2 = 0.89$ at

40 the ATTO pristine tropical rainforest site in central Amazonia). Under polluted conditions, we ob-
41 served strongly enhanced MEK mixing ratios. Overall, the MEK mixing ratios and flux data pre-
42 sented here indicate that both biogenic and anthropogenic sources contribute to its occurrence in
43 the global atmosphere.

44

45 **Keywords:** Methyl ethyl ketone, plant emission, anthropogenic sources, air quality, rainforest, bo-
46 real forests.

47

48 **1. Introduction**

49 Methyl ethyl ketone (C₄H₈O; MEK, also known as 2-butanone, butanone, methyl acetone,
50 butan-2-one, methylpropanone, ethylmethylketone and methylacetone) is an oxygenated volatile
51 organic compound (OVOC). Its occurrence in the atmosphere has been reported over a wide range
52 of environments (Cecinato et al., 2002; Hellén et al., 2004; Ho et al., 2002; Kim et al., 2015;
53 McKinney et al., 2011; Singh et al., 2004) with typical mixing ratios of 0.03 - 4 ppb (Ciccioli and
54 Mannozi, 2007; Kim et al., 2015). Although often being measured alongside other volatile organic
55 compounds (VOCs), atmospheric MEK has received little attention to date. The photochemistry of
56 acetone may serve as an example of how ketones affect the composition and chemistry of the at-
57 mosphere by delivering free radicals to the upper troposphere (Colomb et al., 2006; Finlayson-Pitts
58 and Pitts, 2000; McKeen et al., 1997), and thus increasing the ozone formation potential and alter-
59 ing the oxides of nitrogen (NO_x) regime (Ciccioli and Mannozi, 2007; Folkins et al., 1998; Prather
60 and Jacob, 1997). This understanding may be transferred to MEK, as this molecule is structurally
61 similar to acetone with a comparable absorption spectrum (Martinez et al., 1992). Several studies
62 report that the mixing ratio of MEK in the free troposphere is roughly one quarter of that of acetone
63 (Moore et al., 2012; Singh et al., 2004). However, MEK is about an order of magnitude more
64 reactive than acetone with respect to the hydroxyl radical (OH) (Atkinson, 2000), which makes it
65 a compound of interest in ongoing discussions about the inability to fully account for the reactivity
66 of OH (Nölscher et al., 2016).

67 There are several known but poorly characterized sources of MEK to the atmosphere. Ter-
68 restrial vegetation (Bracho-Nunez et al., 2013; Brilli et al., 2014; Davison et al., 2008; De Gouw

69 et al., 1999; Isidorov et al., 1985; Jardine et al., 2010; Kirstine et al., 1998; König et al., 1995;
70 McKinney et al., 2011; Ruuskanen et al., 2011; Song and Ryu, 2013; Steeghs et al., 2004; Wilkins,
71 1996; Yáñez-Serrano et al., 2015), fungi (Wheatley et al., 1997) and bacteria (Song and Ryu, 2013;
72 Wilkins, 1996) are known to emit MEK. It is also emitted directly by several anthropogenic
73 sources, including man-made biomass burning (Andreae and Merlet, 2001), solvent evaporation
74 (Kim et al., 2015; Legreid et al., 2007) and vehicle exhaust (Bon et al., 2011; Brito et al., 2015; Liu
75 et al., 2015; Verschueren, 1983). In addition, MEK can be formed via the atmospheric oxidation
76 of other compounds (de Gouw et al., 2003; Jenkin et al., 1997; Neier and Strehlke, 2002;
77 Sommariva et al., 2011).

78 Looking in more detail at biogenic sources, MEK emissions have been observed from dif-
79 ferent types of vegetation, including forest canopies (Brilli et al., 2014; Jordan et al., 2009b; Yáñez-
80 Serrano et al., 2015), pasture (Davison et al., 2008; De Gouw et al., 1999; Kirstine et al., 1998) and
81 clover (De Gouw et al., 1999; Kirstine et al., 1998). The MEK production and release mechanisms
82 are manifold, but poorly understood. Studies show higher MEK emissions after cutting and drying
83 of leaves than under no-stress conditions (Davison et al., 2008; De Gouw et al., 1999). Due to the
84 water solubility of MEK in leaves and on surfaces (Sander, 2015), Jardine et al. (2010) suggested
85 MEK emissions to be dependent on evaporation from storage pools in leaves. It has been suggested
86 that MEK takes part in tri-trophic signalling following herbivore attack (Jardine et al., 2010; Song
87 and Ryu, 2013). The roots of plants have also been found to release MEK in root-aphid interactions
88 (Steeghs et al., 2004). Decaying plant tissue may also act as a source of MEK to the atmosphere
89 (Warneke et al., 1999). Furthermore, some studies indicate the importance of MEK emissions by
90 microbes, such as *Brevibacterium linens*, *Bacillus spp* and thermophilic gram-positive actinomy-
91 cetes bacteria (Song and Ryu, 2013; Wilkins, 1996), and fungi such as *Trichoderma spp* (Wheatley
92 et al., 1997).

93 MEK does not only enter the atmosphere via direct emissions, but also results from the
94 atmospheric photooxidation of VOCs such as *n*-butane, 2-butanol, 3-methyl pentane and 2-methyl-
95 1-butene (de Gouw et al., 2003; Jenkin et al., 1997; Neier and Strehlke, 2002; Sommariva et al.,
96 2011). Although butane in the atmosphere comes predominantly from anthropogenic sources
97 (Kesselmeier and Staudt, 1999), some studies have reported emission of *n*-butane from vegetation
98 (Donoso et al., 1996; Greenberg and Zimmerman, 1984; Hellén et al., 2006; König et al., 1995;

99 Zimmerman et al., 1988). The MEK yield from *n*-butane oxidation is ~ 80% (Singh et al., 2004).
100 It is important to note that no mechanistic pathways have been found for atmospheric MEK pro-
101 duction from the dominant biogenic VOCs isoprene, α - and β -pinene and methyl butenol oxidation
102 (Rollins et al., 2009; Singh et al., 2004).

103 In the atmosphere MEK reacts mainly with OH ($k_{OH}=1.15\times 10^{-12}$ cm³ s⁻¹) (Chew and
104 Atkinson, 1996), while reactions with O₃ and NO₃ are very slow during the day and hence negligi-
105 ble (Atkinson and Arey, 2003). MEK has a lifetime of 5.4 days at an OH concentration of 1.6×10^6
106 radicals cm⁻³, whereas isoprene and acetone have lifetimes of 8.2 h and 38 days, respectively, under
107 the same conditions (Grant et al., 2008). The atmospheric degradation of MEK leads to acetalde-
108 hyde and formaldehyde formation. In the presence of NO_x, MEK can lead to peroxyacetyl nitrate
109 (PAN) and ozone formation (Grosjean et al., 2002; Pinho et al., 2005). In the upper troposphere,
110 MEK photolyzes and regenerates OH (Atkinson, 2000; Baeza Romero et al., 2005; De Gouw et
111 al., 1999) as does acetone, potentially increasing ozone formation.

112 Anthropogenic biomass burning leads to significant MEK emissions, of about 2 Tg a⁻¹ glob-
113 ally (Andreae and Merlet, 2001 and unpublished updates; Schauer et al., 2001). Furthermore, about
114 9 Tg a⁻¹ of other C₄ compounds are emitted by biomass burning, which may act as MEK precursors.
115 Another strong source of MEK is biofuel and charcoal combustion, with emissions of ~830 mg kg⁻¹
116 ¹ of dry biomass (compared to an emission rate of ~260 mg kg⁻¹ of dry mass for biomass burning
117 of savannah and grassland vegetation types, Andreae and Merlet, 2001). Despite the fact that bio-
118 mass burning emission rates have been fairly well characterized, vehicular emissions, food cook-
119 ing, industrial activities, cigarette smoke and other anthropogenic sources have not been character-
120 ized. Even though MEK is present in urban atmospheres, there are no observations of MEK emis-
121 sions from vehicles. Most of the urban MEK is released to the atmosphere via evaporation from
122 chemical plants and industrial and household applications, as it is widely used as a solvent (e.g. in
123 glues and as paint thinner). It has a low toxicity, and is not carcinogenic (National Center for
124 Biotechnology, 2015). As its manufacturing has been increasing in the last 10 years, global atmos-
125 pheric mixing ratios have probably increased as well.

126 Here we report recent findings on MEK from six different sites, including biogenic and
127 anthropogenic dominated environments, in order to understand MEK sources in different environ-
128 ments. Our large dataset allows a closer view of this important, almost ubiquitous species in Earth's
129 atmosphere.

130 **2. Sites and Methodology**

131 The field sites compared in our study cover areas from pristine to remote anthropogenically
132 influenced tropical forests, as well as boreal and Mediterranean regions. Measurements were per-
133 formed by proton transfer reaction – mass spectrometry (PTR-MS) and partly complemented by
134 gas chromatography - flame ionization detector (GC-FID) and gas chromatography - mass spec-
135 trometry (GC-MS) analytical techniques (Figure 1, Table 1).

136 Online MEK measurements were performed with quadrupole PTR-MSs (Ionicon Analytic
137 GmbH, Austria, Lindinger et al., 1998) at all sites, except for CYPHEX where a PTR-Time-of-
138 Flight-MS (PTR-ToF-MS, Ionicon Analytic GmbH, Austria, Jordan et al., 2009a) was used. The
139 PTR-MSs were operated at standard conditions (2.2 mbar drift pressure, 600 V drift voltage, 142
140 Td for ATTO and SMEAR-Estonia; 2.0 mbar drift pressure, 550 V drift voltage, 129 Td, for TT34;
141 2.2 mbar drift pressure, 600 V drift voltage, 135 Td for O₃HP; 2.2 mbar drift pressure, 560 V drift
142 voltage, 132 Td for T2 and 2.2 mbar drift pressure, 600 V drift voltage, 137 Td for CYPHEX).

143 Periodic background measurements and weekly humid calibrations were performed at all
144 sites. Gravimetrically prepared multicomponent standards were obtained from Apel & Riemer,
145 USA, for ATTO, TT34, T2 and CYPHEX, and from Ionicon Analytik GmbH, Austria, for O₃HP
146 and SMEAR-Estonia.

147 **2.1. The Amazon Tall Tower Observatory, ATTO: pristine tropical rainforest (Ama-** 148 **zon, Brazil)**

149 The Amazon Tall Tower Observatory (ATTO) site is located in central Amazonia, 150 km
150 NE of Manaus, Brazil (Figure 1) within a pristine primary tropical rainforest. The site is equipped
151 with a tall tower (325 m) and two 80 m towers. One of them (02°08'38.8" S, 58°59'59.5" W) is an
152 80-m walk-up tower, where the trace gas measurements take place. It is surrounded by a forest with
153 a canopy height of approximately 35 m and with at least 417 different tree species among 7293

154 screened trees of ≥ 10 cm diameter at breast height (DBH) in the twelve 1-ha inventoried plots
155 (Andreae et al., 2015). The climate of this site is typical for tropical rainforests with a drier season
156 (July-October) and a wet season (December-April, Nobre et al., 2009).

157 Measurements for this study took place 18 February - 15 March 2014. They were carried
158 out at seven different heights (0.05, 0.5, 4, 24, 53 and 79 m) with the PTR-MS switching sequen-
159 tially between each height in 2 minutes intervals. The inlet lines were made of PTFE, 9.5 mm OD,
160 insulated and heated to 50 °C and had PTFE particle inlet filters. More information about the gra-
161 dient system and PTR-MS operation at ATTO can be found elsewhere (Nölscher et al., 2016;
162 Yáñez-Serrano et al., 2015). The limit of detection (LOD) and uncertainty of the PTR-MS were
163 0.05 ppb and 34% respectively.

164 Additionally, ambient samples for off-line measurements with GC-FID were taken on 11
165 March 2014 from 08:30 to 11:00 LT. They were collected at 24 m using a GSA SG-10-2 personal
166 sampler pump and adsorber tubes (Carbograph 1, Carbograph V 130 mg of Carbograph 1 (90 m²
167 g⁻¹) followed by 130 mg of Carbograph 5 (560 m² g⁻¹)). The size of the Carbograph particles was
168 in the range of 20–40 mesh. Carbograph 1 and 5 were provided by Lara s.r.l. (Rome, Italy). Samples
169 were collected for 20 min with a flow of 167 ml min⁻¹ passing about 3.3 l of ambient air through
170 the adsorbent. Cartridges were transported to the laboratory for analysis by a Perkin Elmer Auto-
171 system XL GC-FID. These samples generally matched the results of the PTR-MS. For details on
172 sampling see Kesselmeier et al., (2002).

173 **2.2. TT34: remote tropical rainforest (Amazon, Brazil)**

174 The ZF2 site is located in the Reserva Biologica do Cuieiras in central Amazonia, 60 km
175 NNW of Manaus (2°35'39.4"S 60°12'33.4"W) within a remote primary tropical rainforest (Figure
176 1). The site is equipped with two towers, TT34 and the K34. The TT34 triangular tower is 40 m
177 high and embedded within the forest with a canopy height of approximately 30 m. The biodiversity
178 of this site is also high and the climate is very similar to that at the ATTO site. More information
179 about the site can be found elsewhere (Karl et al., 2009; Martin et al., 2010).

180 Measurements for this study were made from 1 September 2013 to 20 July 2014 at 41 m,
181 at a fast rate (0.5 Hz) for virtual disjunct eddy covariance (vDEC) flux derivations techniques (Karl

182 et al., 2002; Langford et al., 2009; Rinne et al., 2002). The high-resolution data was further aver-
183 aged to give 30-minute concentration and flux data. Wind vector data were obtained with a sonic
184 anemometer (Gill R3, USA) mounted at the top of the tower close to the PTR-MS inlet. The PTR-
185 MS inlet line was made of PFA (12.7 mm OD) (PFA-T8-062-100, Swagelok), and was insulated
186 and heated to 40 °C inside the air-conditioned cabin. The LOD and uncertainty averaged 30-minute
187 concentrations measurements were 0.18 ppb and 6 % respectively.

188 **2.3. Station for Measuring Ecosystem-Atmosphere Relations, SMEAR-Estonia: re-** 189 **remote hemi-boreal forest (Tartumaa, Estonia)**

190 The Station for Measuring Ecosystem-Atmosphere Relations (SMEAR-Estonia) site is lo-
191 cated in the Järvselja Experimental forest station in Tartumaa, SE Estonia (58°16'N 27°16'E),
192 within a remote hemi-boreal zone, far from major anthropogenic disturbances (Noe et al., 2011,
193 Figure 1). The site is equipped with a tower of 24 m height. The surrounding canopy is about 16-
194 20 m in height and the remote hemi-boreal forest consists of a mixture of tree species, with Norway
195 spruce (*Picea abies*) dominating. The climate is boreal with a growing season of 170-180 days.
196 More information about the site can be found elsewhere (e.g. Bourtsoukidis et al., 2014a; Noe et
197 al., 2011, 2016).

198 The measurements were made between 3 and 17 October 2012. Sampling was done using
199 a dynamic, automated glass enclosure with measurement cycles of 36 seconds. The inlet line (9.5
200 mm OD) was made of glass and was insulated and heated to 70°C. A dynamic exchange enclosure
201 was used to measure emission rates from a Norway spruce branch located in the upper canopy at
202 16 m. While the focus of this study was the quantification of emission rates of MEK from a Norway
203 spruce tree, ambient mixing ratios were derived as well using the box model described in
204 Bourtsoukidis et al., (2014b). The LOD and uncertainty of the PTR-MS were 0.04 ppb and 28%
205 respectively.

206 Furthermore, at SMEAR-Estonia, off-line measurements with a GC-MS were carried out
207 for periods of three days each in June and July 2012, with samples taken every 4 hours at two
208 heights (2 m and 20 m). Samples for GC-MS analysis were also taken from cuvettes enclosing
209 some common plant species at the site (Table 1). In addition, VOC emissions from soil litter were
210 monitored monthly. The air samples were drawn into multi-bed stainless steel cartridges (10.5 cm

211 length, 3 mm inner diameter, Supelco, Bellefonte, PA, USA) filled with Carbotrap C 20/40 mesh
212 (0.2 g), Carbopack C 40/60 mesh (0.1 g) and Carbotrap X 20-40 mesh (0.1 g) adsorbents (Supelco).
213 Even though the site usually experiences low ozone mixing ratios of 10 - 30 ppb (Noe et al., 2012),
214 a catalytic Cu(II) ozone scrubbing system (Sun et al., 2012) was applied. Three constant-flow air
215 sample pumps (1003-SKC, SKC Inc., Huston, TX, USA) and one multisample constant-flow air
216 sample pump (224-PCXR8, SKC Inc., Huston, TX, USA) allowed four samples to be collected at
217 the same time. Each sample took 30 min with a flow of 200 ml min⁻¹ concentrating 6 l of ambient
218 air onto the adsorbent. More information can be found elsewhere (Noe et al., 2012).

219 **2.4. Observatoire de Haute Provence, O₃HP: rural Mediterranean temperate forest** 220 **(Provence, France)**

221 The oak observatory (O₃HP, <https://o3hp.obs-hp.fr>) at the “Observatoire de Haute Pro-
222 vence” (OHP, <http://www.obs-hp.fr/welcome.shtml>), is located within a rural Mediterranean tem-
223 perate forest in the French Mediterranean region, 60 km north of Marseille, the closest large city
224 (43°55'54.0" N 5°42'43.9" E, Figure 1). A 10 m mast was set up inside the oak forest with a canopy
225 height of approximately 5 m. The O₃HP site is dominated by *Quercus pubescens* Willd (75% of
226 trees) and *Acer monspessulanum* L. (25%) forming a sparse canopy, while *Cotinus coggygria* Scop.
227 and other grass species constitute the understorey canopy. The climate at the site is typical Medi-
228 terranean, with dry and hot summers and humid and cool winters. More information about the site
229 can be found elsewhere (Genard-Zielinski et al., 2015; Kalogridis et al., 2014).

230 The measurements took place during 29 May - 12 June 2014 as part of the CANOPEE
231 project (Biosphere-atmosphere exchange of organic compounds: impact of intra-canopy pro-
232 cesses). Ambient measurements were carried out at 2 m (inside the canopy) on consecutive days in
233 intervals of 5 minutes. The 9.5 mm OD Teflon inlet lines were insulated and heated above ambient
234 temperature and had no particle filter. The LOD and uncertainty of the PTR-MS were 0.11 ppb and
235 20% respectively. In addition, light non-methane hydrocarbons (from ethane to hexane) were meas-
236 ured with a GC-FID (Chromatotec, Saint-Antoine, France) in-line with the PTR-MS as described
237 in Zannoni et al. (2016).

238 **2.5. T2: mixed urban and rainforest influenced environment (Amazon, Brazil)**

239 The T2 site is part of a set of experimental sites within the GoAmazon project to study the
240 effect of the pollution plume from the city of Manaus on the otherwise pristine Amazonian atmos-
241 phere. The T2 site is located 8 km downwind, i.e. to the west, of Manaus (3°8'21.12" S, 60°7'53.52"
242 W, Figure 1). Given its location, near Manaus and across the Rio Negro, air mass transport to the
243 sampling site is strongly modulated by a river breeze, alternating between mostly biogenic condi-
244 tions, resulting from the surrounding forest, and the city emissions. The climate is tropical and
245 similar to that at the ATTO and ZF2 sites.

246 The measurements for this study took place between 15 February and 15 November 2014.
247 They were carried out at 12 m above the laboratory container with 30 minute cycles. The inlet line
248 was made of insulated Teflon (9.5 mm OD) without PTFE particle filter. The LOD and uncertainty
249 of the PTR-MS were 0.02 ppb and ~30% respectively.

250

251 **2.6. CYPHEX: mixed marine, rural environment influenced by aged air masses (Cy-** 252 **prus)**

253 The Cyprus Photochemistry Experiment (CYPHEX) campaign took place at a site located
254 in the NW inshore part of Cyprus, in the Paphos region (34°57'50.0" N, 32°22'37.0" E) (Figure 1).
255 The site experiences mixed marine and rural emissions influence. The climate is Mediterranean,
256 warm and dry, and shrubs and small trees dominate the sparse vegetation.

257 The measurements took place during July and August 2014 without a single rain event.
258 Instruments were installed inside containers and connected to a stack inlet that reached up 5 m
259 above the container roofs. Air was drawn through the 8 m stack inlet of 0.5 m with high flow rate
260 (10 l min^{-1}) to minimize wall losses. The subsampling inlet line was made of Teflon (13 mm OD),
261 was insulated and heated to 35 °C, and had a PTFE inlet particle filter. The LOD and uncertainty
262 of the PTR-MS were 0.02 ppb and 11% respectively.

263 **3. Results**

264 **3.1. Sites dominated by biogenic emissions**

265 All the pristine or remote sites studied were characterized by relatively low mixing ratios
266 of nitrogen oxides (NO_x) (< 3 ppb of nitrogen dioxide (NO_2) for O₃HP (Kalogridis et al., 2014;
267 Zannoni et al., 2016), 0.2-0.8 ppb of NO_x for SMEAR-Estonia, and <1 ppb NO_x for the Amazon

268 rainforest (Kuhn et al., 2010)). The diel cycles of MEK at these sites followed a comparable pattern
269 (Figure 2), where MEK mixing ratios were highest in the middle of the day, following the maxima
270 of light and air temperature. The dominant source at these sites was considered to be biogenic.
271 Mixing ratios of MEK correlated well with ambient temperature ($r^2=0.57$ (ATTO), $r^2=0.83$
272 (SMEAR), $r^2=0.47$ (O₃HP)), while it was less well correlated with photosynthetically active radi-
273 ation (PAR) ($r^2= 0.23$ (ATTO), $r^2=0.26$ (SMEAR), $r^2=0.67$ (O₃HP)). This suggests that ambient
274 temperature predominantly influenced MEK emission rates from plants.

275 The vertical observations at ATTO revealed a strong diel variability of the magnitude and
276 vertical distribution of MEK mixing ratios throughout the forest canopy and in the atmosphere
277 above. Figure 3 shows an example of one day (7th of March 2014) hourly vertical profile of MEK
278 from 13:00-15:00 LT, from the ground to the atmosphere, suggesting that the canopy top is the
279 major source of MEK at the site on such days. Similar concentration gradients were found for 83%
280 (for the afternoon hours) and 45% (for the morning hours) of all days of measurements. In addition,
281 MEK mixing ratios decreased significantly beneath the canopy towards the forest floor, possibly
282 due to dry deposition or generally smaller vegetation emissions due to less light and temperature.
283 However, a possible production from the ozonolysis of alkanes or bidirectional plant exchange
284 cannot be ruled out. For a seasonal comparison, Yañez-Serrano et al., 2015 reported 0.43 ppb of
285 MEK for the dry season (September 2013) and 0.13 ppb of MEK for the wet season (February-
286 March 2013) at 38m. Curiously, at 24m, MEK mixing ratios for the wet season were 0.38 ppb, very
287 close to the measured values for this study. Possible differences in canopy structure temperature
288 and solar radiation among years may be the cause for this discrepancy.

289 At the TT34 rainforest site, ecosystem-scale fluxes were directly calculated from the PTR-
290 MS measurements using the method of virtual disjunct eddy covariance (vDEC) (Karl et al., 2001b;
291 Figure 4). The fluxes averaged over the entire 11-month measurement period (covering both parts
292 of dry and wet season) clearly demonstrate an emission of MEK by the rainforest during daytime
293 with the highest emissions around noon, and no emissions during nighttime. In terms of seasonal
294 variation, MEK mixing ratios were observed to be higher during the dry season (September-Octo-
295 ber 2013, ~0.6 ppb) and lower during the wet season (January-April, ~0.2 ppb) (data not shown).

296 Online ambient mixing ratios of MEK, as measured by the PTR-MS in the hemi-boreal
297 forest at the SMEAR-Estonia site during autumn 2012, were on average 0.15 ± 0.04 ppb (range

298 0.09 - 0.25 ppb). These mixing ratios are almost a factor of two lower than ATTO and O₃HP during
299 daytime hours. This difference among boreal forests, with growing season ending in October,
300 broad-leafed tropical (ATTO) and temperate (O₃HP) forests could be partly related to the temper-
301 ature dependence of MEK emissions apparently common among all biogenic sites.

302 The rural Mediterranean temperate forest site at O₃HP differs significantly from the tropical
303 rainforest (ATTO, TT34) or the hemi-boreal forest (SMEAR-Estonia, Figure 1). The trees at O₃HP
304 are predominantly *Quercus pubescens*, a high isoprene emitter (Keenan et al., 2009). At this site,
305 the exchange of air through the forest canopy is enhanced because the canopy is sparse. As shown
306 in Figure 2, ambient MEK mixing ratios measured inside the canopy (2 m) increased with temper-
307 ature in the morning. During the day, increased forest emissions of MEK seemed to balance the
308 rise of the boundary layer depth, resulting in a plateau until sunset. The fluctuation of MEK after
309 sunset may be understood as a result of a ceased source revealing the deposition as it can hardly be
310 explained by gas-phase chemistry or the reduced nocturnal boundary layer height.

311 During the CANOPEE campaign at the O₃HP site, additional GC-FID samples were taken
312 at 2 m, allowing measurements of several anthropogenic light hydrocarbons, including n-butane.
313 This sampling was performed in parallel to the PTR-MS measurements. All samples contained n-
314 butane, which typically has an anthropogenic origin with an average mixing ratio of 0.071±0.09
315 (much lower mixing ratios than MEK), indicating there is no significant source of n-butane nearby.
316 Hence, MEK at the O₃HP site could not be related to the atmospheric oxidation of n-butane. Fur-
317 thermore, the absence of a correlation with other anthropogenic tracers lets us conclude that MEK
318 at this site was predominantly of biogenic origin.

319 The measurements obtained by PTR-MS at the presented sites dominated by biogenic emis-
320 sions were occasionally confirmed by GC-FID and GC-MS, which are compound selective. At
321 ATTO the same range of MEK mixing ratios for the same hour of the day and height for the GC-
322 FID and the PTR-MS measurements was found, indicating that the PTR-MS signal was only or at
323 least dominated by MEK. To identify sources, canopy measurements at SMEAR-Estonia were
324 complemented by emission measurements using cuvettes with GC-MS identification. Common
325 hemi-boreal forest species, such as *Quercus robur*, *Tilia cordata*, *Sorbus aucuparia*, *Betula pu-*
326 *bescens* and *Picea abies*, were screened for VOC emissions. The highest emissions of MEK were
327 found from *Tilia cordata* and *Picea abies* (Table 2). The data are consistent with those reported by

328 (Bourtsoukidis et al., 2014b) who measured an emission rate of MEK of $2.6 \pm 2.2 \text{ ng g}_{(\text{dw needle})}^{-1} \text{ s}^{-1}$
329 from *Picea abies* using PTR-MS. The GC-MS technique obtained a very similar value of 2.3 ng
330 $\text{g}_{(\text{dw needle})}^{-1} \text{ s}^{-1}$. MEK emissions from *Picea abies* were relatively small compared to other VOCs
331 emitted by the same plant species such as total monoterpenes and acetone which reached 93.2 and
332 $27.6 \text{ ng g}_{(\text{dw needle})}^{-1} \text{ s}^{-1}$, respectively (Bourtsoukidis et al., 2014b). In addition to plant sources, the
333 emissions rates of MEK from leaf litter were quantified using a GC-MS. MEK litter emissions,
334 with a monthly average of $7 \mu\text{g m}^{-2} \text{ h}^{-1}$, were of comparable magnitude to the emissions rates of
335 MEK from the screened hemi-boreal forest tree species, including *Quercus robur* or *Sorbus aucu-*
336 *paria*, which emitted 8 - 9 $\mu\text{g m}^{-2} \text{ h}^{-1}$ of MEK (Table 2).

337 **3.2. Anthropogenically influenced sites**

338 Anthropogenically influenced sites are characterized by air masses that have passed over
339 polluted cities or industrialised regions. This air typically has elevated mixing ratios of NO_x , other
340 products of fossil fuel combustion such as aromatic compounds, carbon monoxide (CO), and higher
341 aerosol loading. Regional biomass burning plumes can also be a source of anthropogenic input to
342 air masses and are here considered as anthropogenic.

343 The T2 data set was sorted for polluted periods (air masses loaded with CO, black carbon,
344 high aerosol loading, aromatic compounds) and non-polluted periods. Periods with CO higher than
345 130 ppb during the tropical wet season and higher than 160 ppb during the dry season were con-
346 sidered polluted. As shown in Figure 5, MEK mixing ratios strongly increase with pollution. The
347 T2 site in Brazil is located on the bank of the Rio Negro and is affected by both, the tropical rain-
348 forest (biogenic) and the megacity of Manaus (anthropogenic). The location of the T2 site down-
349 wind of Manaus and upwind of the rainforest minimizes the biogenic influence. MEK mixing ratios
350 were generally lower for the clean conditions at T2 than mixing ratios found at ATTO or TT34
351 (Figure 2 and 5). Nevertheless, the mixing ratios of MEK during polluted conditions (0.7 ± 0.33
352 ppb during dry season at 13:00 LT) reached or even exceeded those at the other tropical rainforest
353 sites (0.32 ± 0.13 ppb at 13:00 LT for ATTO and 0.45 ± 0.28 ppb at TT34; Figure 2 and 5). Pre-
354 sumably, anthropogenically affected air as transported across the Rio Negro from the city of Ma-
355 naus (~2 million inhabitants; IBGE, 2014) generated a plume with a mixture of anthropogenic
356 MEK directly emitted in Manaus or MEK formed by oxidation of e.g. *n*-butane during transport.
357 The general trend observed in Figure 6 is an increase of MEK mixing ratios when easterly winds

358 came from Manaus (located to the East of T2). However, there were times when winds blew from
359 the North, and incident air masses passed through large rainforest areas, in which TT34 is included,
360 these air masses then crossed the river and arrived at the site). During these times, when air masses
361 were mostly dominated by biogenic emissions, MEK enhancement reached on average 0.2 ppb.

362 Mixing ratios of MEK at T2 were found to be significantly enhanced during polluted con-
363 ditions for both dry and wet season (Figure 7). The relative enhancement within polluted periods
364 at 13:00 LT ranged around a factor of 1.5 for the wet season and of 1.8 for the dry season. During
365 the dry season, the day-to-day variability was more intense, as reflected by the standard deviations
366 which increased by 360 % for the clean conditions and 410 % for the polluted conditions relative
367 to the wet season clean and polluted values, respectively. This may indicate a difference in the
368 sources and sinks regulating MEK mixing ratios among the different seasons. Examples of this
369 difference could be an increase of MEK due to biomass burning, more abundant during the dry
370 season, or changes in the deposition rates due to changes in rain frequency.

371 The CYPHEX campaign took place at Ineia, North-West Cyprus, at a location that has very
372 little significant vegetation nearby. The air masses that pass through the site are either from Western
373 Europe passing across France and Spain and then the Mediterranean Sea, or Eastern Europe (e.g.
374 Turkey, Greece). During the CYPHEX campaign, the hourly median MEK mixing ratios did not
375 show any distinct diel cycle or relations to temperature or net radiation (Figure 5) strongly suggest-
376 ing that no significant local sources were present. Furthermore, backward air mass trajectories, as
377 calculated by the HYSPLIT model (NOAA Air Resources Laboratory, USA, Stein et al., 2015)
378 (Figure 8), can be used to delineate times when Cyprus was affected by easterly and westerly flow.
379 These trajectories were started at 650 m height with the ensemble mode. The periods (east, west)
380 were chosen on the basis of the FLEXPART model. Further information can be found in Derstroff
381 et al., in preparation. On average, easterly air masses contained 0.13 ± 0.03 ppb whereas westerly
382 masses contained 0.08 ± 0.02 ppb. This difference can be due to differences in source strength, the
383 greater duration of boundary layer transport from the west and hence marine uptake, or to photo-
384 chemical loss during transport.

385 **3.3. Compilation of measurement data**

386 In order to investigate the origin and characteristics of MEK in the atmosphere, we calcu-
387 lated the correlation coefficient (r^2) between the mixing ratios of MEK and other co-measured VOC
388 species for each site (Table 3). This coefficient is the ratio of the variability of the MEK mixing
389 ratios over the variability of the other VOC mixing ratios available for each site. The correlation
390 between MEK and other VOC helps to identify possible similarities such as common sources,
391 sinks, chemical mechanisms, or transport processes. However, this comparison does not neces-
392 sarily claim links between the various compounds. Acetone, acetaldehyde, monoterpenes, isoprene,
393 isoprene oxidation products and methanol are regarded as being of biogenic origin especially in
394 forested areas (Kesselmeier and Staudt, 1999; Laothawornkitkul et al., 2009). Compounds such as
395 benzene, toluene, xylene and acetonitrile are considered as typical anthropogenic tracers (Andreae
396 and Merlet, 2001; Finlayson-Pitts et al., 1997).

397 In general, biogenic sites, namely ATTO, SMEAR-Estonia, and O₃HP, showed relatively
398 high correlations between MEK and almost all biogenic VOCs ($r^2 > 0.5$). Exceptions are the r^2 of
399 isoprene, monoterpene and isoprene oxidation products for O₃HP. For instance, the highest corre-
400 lation coefficient (r^2) was found for MEK and acetone at the SMEAR site ($r^2 = 0.97$). A high cor-
401 relation coefficient could indicate that the atmospheric processes governing acetone and MEK
402 abundance are likely very similar (Zhou and Mopper, 1993). In SMEAR-Estonia overall high cor-
403 relations were found between MEK and the oxygenated compounds, acetone, acetaldehyde and
404 methanol, as well as with monoterpenes and isoprene. At ATTO, correlations were only slightly
405 lower. The correlation coefficient (r^2) calculated for the O₃HP observations were generally lower
406 than for ATTO and SMEAR-Estonia, further influenced by the higher turbulent mixing due to
407 sparser vegetation, leading to fastest transport to the atmosphere. Nevertheless, the good correla-
408 tions of MEK with typical biogenically emitted compounds, such as isoprene, isoprene oxidation
409 products, monoterpenes, methanol and acetone, corroborated the biogenic origin of MEK emis-
410 sions at the biogenic sites.

411 At the anthropogenically influenced sites, T2 and CYPHEX, correlation coefficient (r^2) for
412 the biogenic compounds were generally lower, apart from the r^2 (0.64 and 0.45, respectively) be-
413 tween MEK and acetone. It is important to note that although T2 is a mixed anthropogenic and
414 biogenic site, the correlation coefficient (r^2) for MEK and acetone was high, but very low for the
415 rest of the biogenic compounds. For the anthropogenic compounds, T2 had an r^2 of 0.27 for MEK

416 and acetonitrile and MEK and xylene. Furthermore, the data from the Cyprus site showed poor
417 correlation of MEK with any biogenic compound, but a correlation coefficient of $r^2 = 0.58$ for
418 MEK and toluene, an anthropogenic tracer.

419 **4. Discussion**

420 **4.1. PTR-MS measurements**

421

422 Most of the measurements in this study were performed with a quadrupole PTR-MS, a
423 technique that monitors selected VOC ions, online and with fast time response. A disadvantage is
424 the separation by masses with a mass resolution of only 1 amu. For some masses, several com-
425 pounds and/or compound-fragments may be detected as one signal. The quadrupole PTR-MS sig-
426 nal at m/z 73 is attributed to MEK, but may have contributing signals of water clusters (de Gouw
427 et al., 2007), butanal (Inomata et al., 2010; McKinney et al., 2011; Slowik et al., 2010; Warneke et
428 al., 2007), acrylic acid (de Gouw et al., 2003), 2-methyl propanal (Baraldi et al., 1999; Jardine et
429 al., 2010), and methyl glyoxal (Holzinger et al., 2007; Jordan et al., 2009b). We have tried to take
430 into account possible interferences by using different analytical techniques and supplementary in-
431 formation. At the SMEAR-Estonia site, the accompanying GC-MS observations validated the sig-
432 nal for MEK. Additionally, the GC-FID samples taken at ATTO corroborated the signal for MEK
433 at this site. Nevertheless, we try to give a short overview below about the interferences of other
434 trace gases with the PTR-MS identification of MEK.

435 Methyl glyoxal is a likely contributor to the observed signal at the PTR-MS protonated
436 mass m/z 73, especially in areas where there are high levels of isoprene. It is formed following the
437 oxidation of methyl vinyl ketone and methacrolein, which are both isoprene oxidation products
438 (Calvert and Madronich, 1987; Lee et al., 2006). Supported by GC-FID measurements and rela-
439 tively low isoprene levels during the wet season (Yáñez-Serrano et al., 2015), we can assume that
440 the contribution of methyl glyoxal to this mass was insignificant at the rainforest sites (ATTO and
441 TT34). Despite the high isoprene emissions at O₃HP, the correlation between MEK and the iso-
442 prene oxidation products was low ($r^2 = 0.41$), suggesting that methyl glyoxal did not significantly
443 contribute to the signal at m/z 73. During the CYPHEX campaign the PTR-ToF-MS could unam-
444 biguously distinguish between MEK and methyl glyoxal (at 73.0648 amu and 73.0284 amu respec-
445 tively).

446 Even though a contribution of butanal to m/z 73 of up to 65% (Lindinger et al., 1998) and
447 20% (Williams et al., 2001) has been reported previously, most butanal fragments on m/z 57 (Ion-
448 icon Analytic GmbH). Acrylic acid, a marine compound (Liu et al., 2016) that may interfere at m/z
449 73, was probably not of relevance at sites under biogenic influence. However, in the case of an-
450 thropogenically influenced sites, such as T2, interferences may have been of relevance. Karl et al.
451 (2007) and Ciccioli et al., (2014) measured tropical biomass burning emissions and found that m/z
452 73 is comprised of 74% MEK and 23% 2-methyl propanal (73.1057 amu). Even though none of
453 the sites presented in this study was severely influenced by biomass burning, we cannot completely
454 rule out a possible direct emission of 2-methyl propanal by plants, which is of lower magnitude
455 than from biomass burning (Hafner et al., 2013; Jardine et al., 2010; Karl et al., 2005a). Due to the
456 standard operation conditions of the PTR-MS under our measurement conditions, we neglected
457 water clusters as they are regarded to be insignificant (McKinney et al., 2011; Yáñez-Serrano et
458 al., 2015). Summarizing these issues, we note that several studies have concluded m/z 73 to origi-
459 nate from MEK only (Bourtsoukidis et al., 2014b; Crutzen et al., 2000; De Gouw et al., 1999, 2000;
460 Holzinger et al., 2000; Karl et al., 2001b, 2005a; Kim et al., 2015; Millet et al., 2015; Steeghs et
461 al., 2004). Based on these considerations and the similarity of magnitudes measured by the PTR-
462 MS as compared with the GC results, we assume m/z 73 is representative of the atmospheric MEK
463 present.

464 **4.2. The biogenic MEK**

465 The data obtained at the biologically influenced sites demonstrated that MEK was emitted
466 by vegetation. This is clearly supported by the canopy-scale net flux observations of MEK at the
467 TT34 rainforest site (Figure 4) as well as the diel cycles of the mixing ratios at the other biogeni-
468 cally influenced sites (Figure 2). Furthermore, the leaf-level cuvette measurements at SMEAR-
469 Estonia also corroborated the MEK emission by vegetation. In addition, a contribution by other
470 biogenic sources such as dead and decaying plant matter was also observed at SMEAR-Estonia to
471 be of similar magnitude to boreal plant species emissions, and indicating a source from plant litter,
472 in accordance with the results from Warneke et al., (1999) that measured MEK emission from the
473 abiotic processes of plant decaying matter. This is not the case for the tropical sites where vertical
474 profiles show that canopy emissions dominate.

475 High correlation coefficients suggested strong relations between the emission processes for
476 MEK and other biogenic compounds (Table 3). A similar approach has been used previously by
477 Goldstein and Schade (2000) to unveil the sources of acetone. Similarly, Davison et al. (2008)
478 found a high correlation coefficient between MEK and acetone of $r^2=0.87$ and a relatively poor
479 correlation between MEK and monoterpenes ($r^2=0.54$). They surmised that good correlations indi-
480 cated a common origin for biogenically emitted compounds. Furthermore, a resemblance of the
481 pattern of acetone and MEK has been reported for the ATTO site before (Yáñez-Serrano et al.,
482 2015). In our study, we found high correlation coefficients (r^2) for MEK with acetone and MEK
483 with temperature, and lower r^2 for MEK and compounds such as isoprene and monoterpenes for
484 all the biogenic sites (Table 3). This could indicate that MEK forest emissions are more related to
485 processes resembling acetone emissions and temperature dependent processes, rather than light and
486 temperature dependent emission mechanisms, as for isoprene and monoterpenes (Jardine et al.,
487 2015; Kesselmeier and Staudt, 1999).

488 Plant physiological production pathways have been reported for MEK formation. MEK can
489 be formed, similarly to acetone, as a by-product of a cyanohydrin lyase reaction during cyanogen-
490 esis (Fall, 2003; Vetter, 2000). This chemical defence pathway was also identified in clover by
491 Kirstine et al. (1998) and de Gouw et al. (1999) as a result of mechanical stress, and can be of
492 special importance for tropical rainforests (Miller et al., 2006). On the other hand, in places such
493 as SMEAR-Estonia, dominating plant species are not cyanogenic, and other processes for MEK
494 formation are probably more dominant. In pine trees, acetone is produced from light-dependent
495 and –independent processes that can be associated with the decarboxylation of acetoacetate occur-
496 ring in microorganisms and animals (Fall, 2003), oxidation of fatty acids leading to ketone emis-
497 sions (Niinemets et al., 2014), from pyruvic acid leading to acetyl-CoA (Kesselmeier and Staudt,
498 1999), or produced from uncharacterized biochemical reactions (Fall, 2003). Such processes could
499 also be related to MEK emissions.

500 Even though extensive laboratory measurements are needed to identify the dominant plant
501 process or processes responsible for MEK emission, this study demonstrated the role that temper-
502 ature can exert on such emissions. Hence, forests around the world may act as very different sources
503 for atmospheric MEK. This can be seen for boreal forests (SMEAR-Estonia), with distinctly lower

504 temperatures, where MEK levels were significantly lower. However, other factors must be consid-
505 ered (Schade et al., 2011), such as Leaf Area Index (LAI) and plant species composition, as well
506 as the environmental factors, water availability and mechanical stress, the latter having already
507 been observed by de Gouw et al. (1999) to act as a driver for MEK emissions.

508 Due to its relatively long atmospheric lifetime (~5 days for the reaction with OH; Grant et
509 al., 2008), MEK is expected to accumulate in the atmosphere until removal. Hence, atmospheric
510 mixing ratios can reflect seasonality and changes in dominating sources, affected by radiation,
511 temperature and phenology, from more biogenic dominance during the wet season to transport
512 phenomena and oxidation processes of primarily emitted compounds from regional biomass burn-
513 ing, as it has been seen in 2013 at the ATTO site (Yáñez-Serrano et al., 2015). Moreover, a possible
514 production from certain terpenes through ozonolysis cannot be excluded (Holzinger et al., 2005).
515 Additionally, the canopy structure seems to be important for air mixing and transport, as seen for
516 the O₃HP site with an apparently faster mixing due to sparser vegetation and consequent dampen-
517 ing of the amplitude of the diel cycle. Furthermore, due to its oxygenated nature, partitioning to
518 and from aqueous surfaces is likely, including deposition and surface reactions. Its high water sol-
519 ubility might allow dissolution within leaf water (Sander, 2015) triggering bidirectional exchange
520 of MEK (Karl et al., 2005b; McKinney et al., 2011; Niinemets et al., 2014). Due to its high solu-
521 bility in water and its relatively long lifetime, MEK could potentially influence gas-aqueous reac-
522 tions on aerosol surfaces (Nozière, 2005). This has been shown indirectly by the production of
523 methyl glyoxal after its oxidation by OH, having implications for the formation of organics in the
524 aerosol aqueous phase (Rodigast et al., 2015).

525 **4.3. The anthropogenic MEK**

526 A clear difference could be observed between the anthropogenic and biogenic influenced
527 sites presented in this study. The T2 site represented a site with mixed influence by urban area and
528 tropical rainforest. Affected by anthropogenic and biogenic sources ambient mixing ratios of MEK
529 were higher than at the pristine ATTO rainforest site. Polluted episodes (from the Manaus plume)
530 with an increase of MEK could be distinguished for both, the wet and the dry season, suggesting a
531 short range transport of air masses. On the other hand, when the wind is blowing from the North,
532 MEK mixing ratios were also present, showing an influence from biogenic forest emissions (Figure
533 7), thus having a mix of biogenic and anthropogenic influence at the T2 site. A strong seasonality

534 of MEK mixing ratios at T2 reflected biomass burning as a common occurrence in the Amazon
535 region during the dry season (Artaxo et al., 2013). In addition to MEK, a higher contribution of
536 butanal affecting m/z 73 (Inomata et al., 2010; Karl et al., 2007) might be possible, although MEK
537 has been reported to have a much higher emission factor (range from 0.17 to 0.83) than butanal
538 (range from 0.04 to 0.21) for biomass burning (Andreae and Merlet, 2001).

539 We regarded CYPHEX as an anthropogenically influenced site with weak or no apparent
540 direct sources, but affected by anthropogenic air masses after long range transport over marine
541 areas. Losses by transport over the sea and chemical decomposition led to the lowest averaged
542 MEK mixing ratios of all compared sites. Correlation coefficients (r^2) of MEK with the biogenic
543 tracers were relatively poor for T2 and CYPHEX. However, correlations were also poor for the
544 anthropogenic tracers, although slightly better than at the biogenic sites. MEK showed highest cor-
545 relation with acetone, indicating similar sources and fate in air mixing and chemistry processes.
546 MEK transported over a long distances (10 days) is lost by photochemical aging or deposition as
547 evidenced by the lowest values reported from CYPHEX. This is despite known secondary photo-
548 chemical sources, i.e. *n*-butane oxidation (Katzenstein et al., 2003; Kwok et al., 1996). Interest-
549 ingly, even under polluted conditions, MEK did not correlate with aromatic compounds, except
550 during CYPHEX, although this correlation deteriorated in the more aged westerly air masses. This
551 can only be understood as a result of a very complex mixture of anthropogenic sources of MEK
552 which vary from direct emission by industry (Legreid et al., 2007), gasoline combustion
553 (Verschueren, 1983), biomass burning (Andreae and Merlet, 2001) and vehicular emissions (Brito
554 et al., 2015). Furthermore, chemical processing during transport may contribute, such as oxidation
555 of *n*-butane, however, the longer transport times during CYPHEX from the west corresponded to
556 lower values.

557 **5. Remarks and conclusions**

558 The comparison of MEK mixing ratios in different parts of the world is necessary in order
559 to understand how this ubiquitous compound occurs and behaves in the atmosphere. Summarizing,
560 Table 4 aims to provide a numerical comparison of MEK mixing ratios reported around the globe.
561 While MEK mixing ratios in our study are relatively constant, MEK has been measured in many
562 different ecosystems ranging from 0.073 ppb to up to 4 ppb. Therefore, it is important to consider
563 the variability of this compound as MEK can lead to PAN and ozone formation in the atmosphere

564 (Pinho et al., 2005). Photochemical odd-hydrogen production in the upper troposphere (Atkinson,
565 2000; Baeza Romero et al., 2005; De Gouw et al., 1999) can further enhance the MEK ozone
566 forming potential (Folkins et al., 1998; Prather and Jacob, 1997).. Of the widely used atmospheric
567 chemistry models, only GEOS-Chem explicitly computes MEK but only with regard to anthropo-
568 genic origin. On the basis of the data presented here from forest sites, it is necessary for atmospheric
569 chemistry models to also include biogenic MEK emissions to better estimate its effects on the
570 environment. Sites under biogenic influence showed marked diel variability, matching biogenic
571 VOC emissions and temperature. Structural forest features seem to affect turbulent mixing and
572 diluting of trace gases like MEK, as in the case of O₃HP with patchy vegetation. MEK seemed to
573 be produced in plants in a similar fashion to acetone, likely released during mechanical stress.
574 Possible pathways for productions in plants are oxidation of fatty acids, cyanogenesis, production
575 from pyruvic acid leading to Acetyl-CoA, light-dependent and –independent processes that can be
576 associated with the decarboxylation of acetoacetate occurring in microorganisms and animals.

577 This study presents the first compilation and comparison of ambient measurements of MEK
578 at different sites. MEK patterns and mixing ratios differ around the globe depending on sources
579 and transport. Vegetation and litter have been identified as sources of MEK and magnitude of
580 sources varied among the tropical rainforest, the Mediterranean temperate forest and the hemi bo-
581 real forest following a likely temperature dependence. However, via different filtering methodolo-
582 gies (CO filtering and backward trajectories), the anthropogenic input from polluted regions, such
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915 **8. Tables**

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Table 1: Measurement sites, site environment, sampling dates, methods used and sampling heights.

Site	Characteristics	Canopy height	Time of Sampling	Technique	Type of measurement	Measurement heights
ATTO (Brazil)	Pristine tropical rainforest	35 m dense veg.	Feb/Mar 2014	PTR-MS GC-FID (samples for off-line analysis, collected volume=3.34 L)	Ambient	0.05, 0.5, 4, 24, 38, 53, 79 m 24 m
TT34 (Brazil)	Remote tropical rainforest	30 m dense veg.	Sep 2013 – Jul 2014	PTR-MS	Ambient, including canopy-scale fluxes	41 m
SMEAR (Estonia)	Remote hemi-boreal forest	16-20 m dense veg.	Jun, Jul, Oct 2012 Oct 2012	GC-MS (samples for off-line analysis, collected volume=6 L) PTR-MS	Ambient, plant, soil enclosure Ambient, plant enclosure	2, 20 m 16 m
O ₃ HP (France)	Rural temperate forest	5 m sparse veg.	May-Jun 2014	PTR-MS	Ambient	2 m
T2 (Brazil)	Mixed urban and rainforest influenced environment	Influence from veg. nearby	Feb-April 2014 July-Oct 2014	PTR-MS	Ambient	14 m
CYPHEX (Cyprus)	Mixed marine, rural environment influenced by aged air masses	None, on top of a hill	Jul-Aug 2014	PTR-TOF-MS	Ambient	8 m

Table 2: Emission rates of MEK for typical hemi-boreal plant species and soil litter measured by GC-MS technique at the SMEAR site.

Plant species and soil cuvettes	Mean $\mu\text{g m}^{-2} \text{h}^{-1}$	Standard deviation $\mu\text{g m}^{-2} \text{h}^{-1}$	Standard error $\mu\text{g m}^{-2} \text{h}^{-1}$	Median $\mu\text{g m}^{-2} \text{h}^{-1}$	Number of data points for statistics
<i>Quercus robur</i>	8.12	-	-	-	1
<i>Tilia cordata</i>	12.93	4.89	3.46	12.93	3
<i>Sorbus aucuparia</i>	9.08	-	-	-	1
<i>Betula pubescens</i>	9.36	5.10	2.94	8.21	3
<i>Picea abies</i>	13.76	5.05	2.91	15.51	3
Leaf litter	7.00	3.37	2.11	6.58	24

Table 3: Correlation coefficients (r^2) of MEK with other co-measured VOC at the sites investigated. The correlations above 0.5 are colour coded with warmest colours for highest correlation coefficients. Correlation coefficients (r^2) for the TT34 site in Amazonia are missing due to lack of data.

r^2		Biogenic tracers					Anthropogenic tracers				
		Acetone	Acetaldehyde	Monoterpenes	Isoprene oxidation products	Methanol	Isoprene	Acetonitrile	Benzene	Toluene	Xylene
	TT34	-	-	-	-	-	-	-	-	-	-
	Biogenic sites										
	SMEAR	0.97	0.89	0.72	-	0.90	0.84	-	-	-	-
	ATTO	0.89	0.62	0.75	0.75	0.51	0.77	0.49	0.07	0.27	0.04
	O₃PH	0.61	0.62	0.12	0.41	0.57	0.14	0.19	0.03	0.15	0.00
	Anthropogenic sites										
	T2	0.64	0.21	-	0.41	0.27	0.06	0.27	0.11	0.07	0.27
	CYPHEX	0.45	0.42	0.07	0.10	0.25	0.08	0.00	0.58	0.09	-

Table 3: Literature compilation of MEK mixing ratios measurements in different ecosystems around the globe from a wide range of atmospheric environments.

Reference	MEK mixing ratio in ppb	Type of environment	Reported data	Height from surface	Time of measurement
Zhou and Mopper, 1993	0.03	Caribbean ocean	Cruise data, total average	10m	October 1988
Ho et al., 2002	0.14	Hong Kong urban centre	annual average	6m	April 2000-April 2001
Cecinatto et al., 2002	0.15	Algeria rural residential site	Daytime average	Not specified	January 2001
Cecinatto et al., 2002	0.23	Algeria rural residential site	Daytime average	Not specified	August 2001
Feng et al., 2004	2.11	China urban site	Evening time	Not specified	August–September 2002
Hellen et al., 2004	0.073	Finnish boreal forest site	Diel average	1 m	March-April 2003
Sighn et al., 2004	0.12	Tropospheric pacific ocean	Non specified	0-2km	February-April 2001
Legreid et al., 2007	0.2*	Swiss urban site	Annual average	8 m	Throughout 2005
Davison et al., 2007	0.8*	Swiss permanent grassland site	Daytime average	1.2 m	June 2005

Grant et al., 2008	~ 0.5	Senegalese rural site	Daytime average	6 m	September 2006
Jordan et al., 2009	0.13	USA mixed rural site	Interannual median from 2005-2008	12 m	From 2005 to 2008
Kim et al., 2015	~4	South Korea semi rural site	Diel average	15 m	June 2013
Yáñez-Serrano et al., 2015	0.43	Brazilian tropical rainforest site (ATTO)	Daytime average	38 m	September 2013
Yáñez-Serrano et al., 2015	0.13	Brazilian tropical rainforest site (ATTO)	Daytime average	38 m	February-March 2013
This study	0.39	Brazilian tropical rainforest site (ATTO)	Daytime average	38 m	February-March 2014
This study	0.39	Brazilian tropical rainforest site (TT34)	Daytime average	41 m	September 2013- July 2014
This study	0.19	Hemi boreal forest site (SMEAR-Estonia)	Daytime average	16 m	October 2012
This study	0.30	French Mediterranean forest (O ₃ HP)	Daytime average	2 m	May-June 2014
This study	0.13	Brazilian mixed Tropical rainforest site(T2)	Daytime average	14 m	February - October 2014
This study	0.11	Cyprian rural site (CYPHEX)	Daytime average	8 m	July -August 2014

*average from different seasons reported. **average of different conditions cutting, a day after cutting removing hay and a day after removing hay.

9. Figures

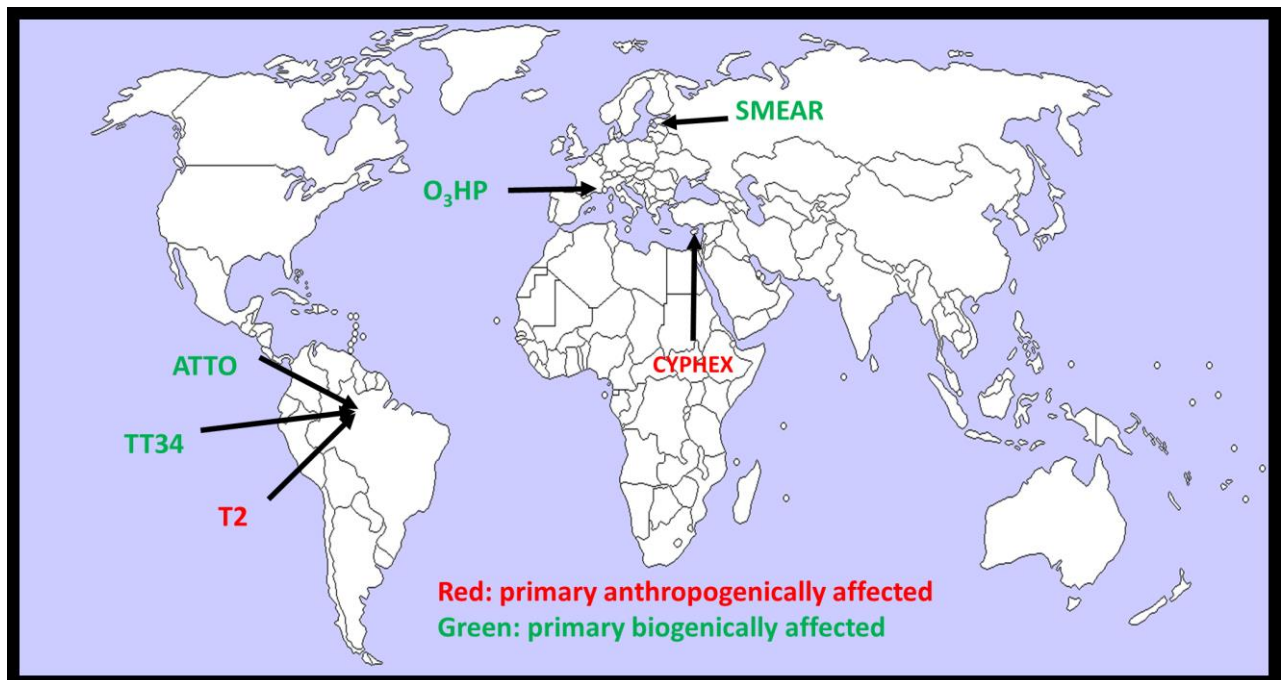


Figure 1: World map showing the location of the different sites. The names are colour coded depending on whether they have primarily biogenic influence (green) or a primarily anthropogenic influence (red). Wikimedia Foundation, 2016.

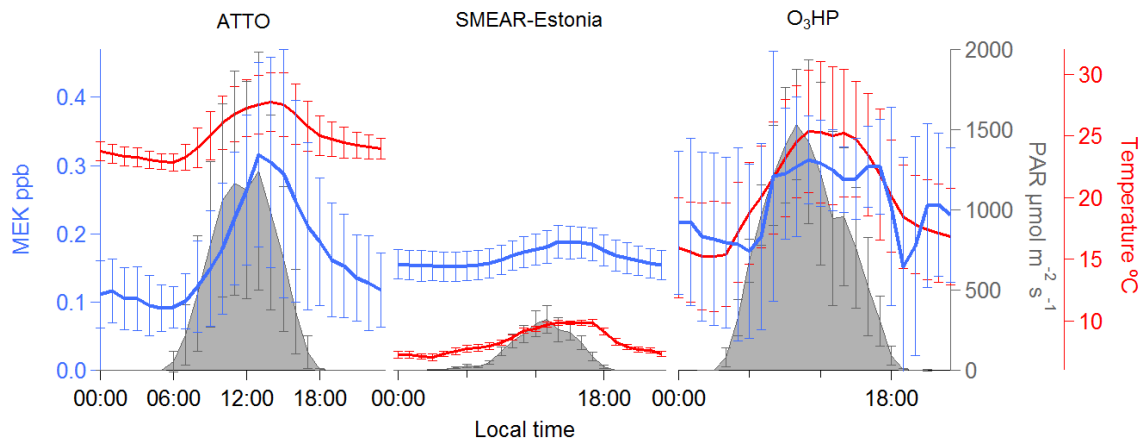


Figure 2: Hourly average diel cycles of MEK at the ATTO (left), SMEAR-Estonia (middle) and O₃HP (right) sites, for the period of measurements (wet season 2014 for ATTO at 38 m October 2014 for SMEAR-Estonia at 16 m, and May and June 2014 for O₃HP at 2 m). Hourly mean diel cycles of temperature and PAR are also shown in red and grey, respectively. Error bars represent the standard deviations.

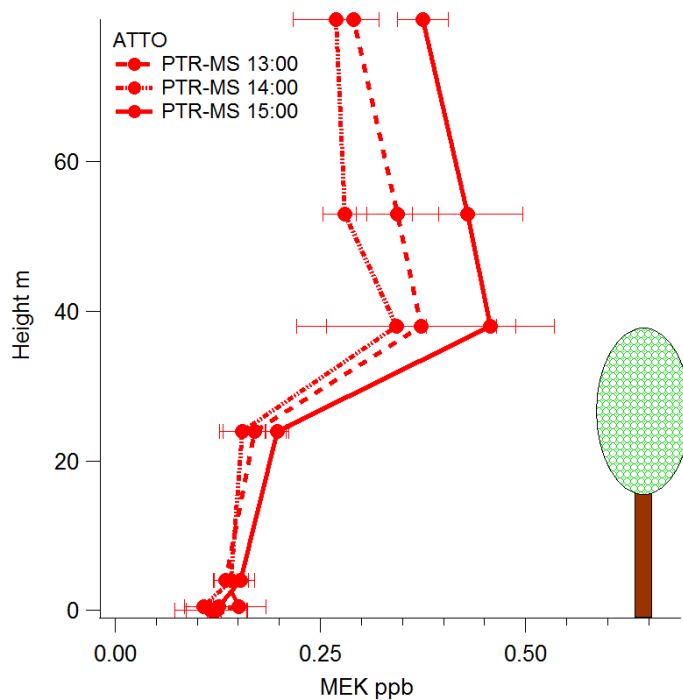


Figure 3: Hourly average vertical profiles of MEK mixing ratios at ATTO for the 7th of March 2014 for 13:00 LT (dashed lines), 14:00 LT (dotted and dashed lines) and 15:00 LT (thick lines). Error bars of vertical profiles are the standard deviations.

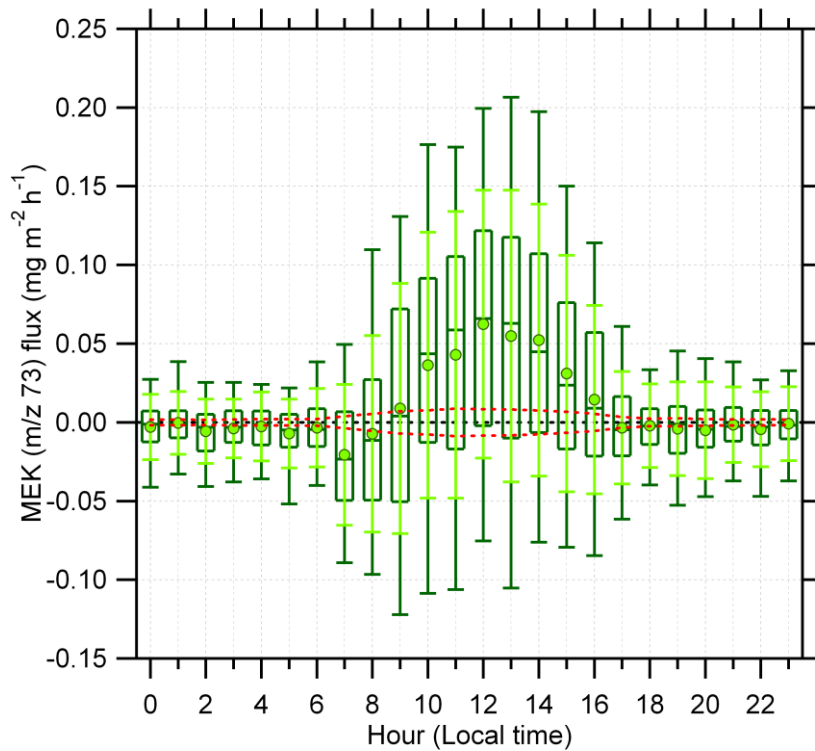


Figure 4: Hourly average MEK fluxes at the TT34 tower for the period Sept 2013 - July 2014. The light green circles represent means and associated error bars are one standard deviation. The central line of the box plots (dark green) indicates the median, bottom and top lines the 25th and 75th percentile respectively and whiskers are the 5th and 95th percentiles. Red dashed lines indicate the propagated limit of detection calculated according to the method outlined by (Langford et al., 2015).

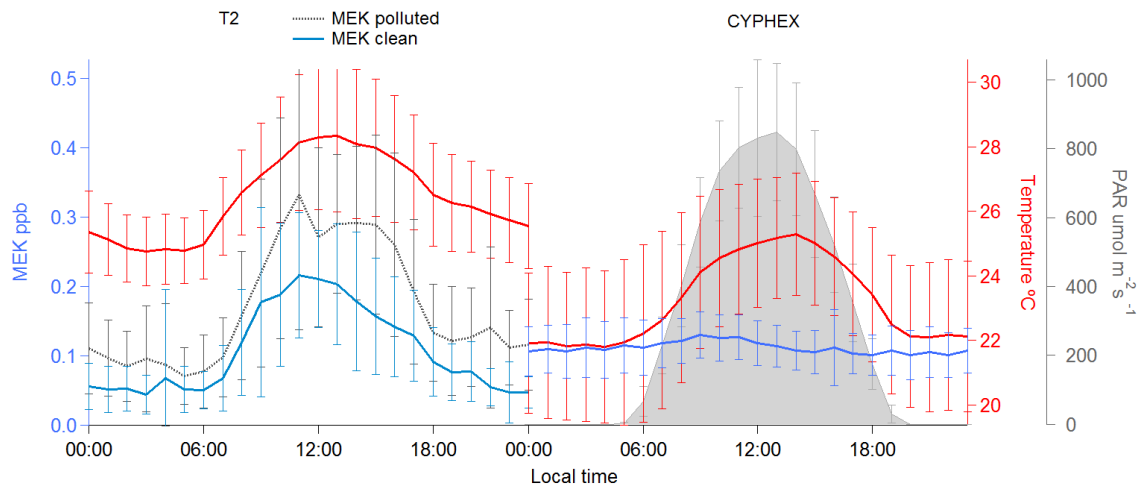


Figure 5: Hourly average diel cycles of MEK at the T2 (left) and CYPHEX (right) sites, for the period of measurements (wet season 2014 for T2 at 14 m, July and August 2014 for CYPHEX at 12 m). For T2 a separation between polluted (dotted black line) and clean (thick blue line) air masses was done. Hourly mean diel cycles of temperature and PAR are also shown in red and grey, respectively. Error bars represent the standard deviations.

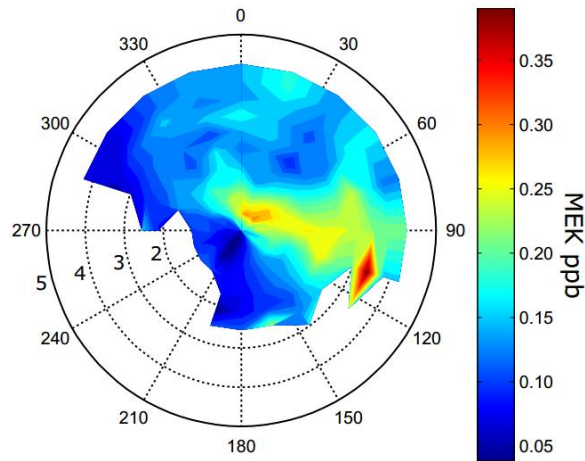


Figure 6: Polar surface plot for average MEK mixing ratios at a given wind direction (angle, $1-5 \text{ m s}^{-1}$) and wind speed (radius).

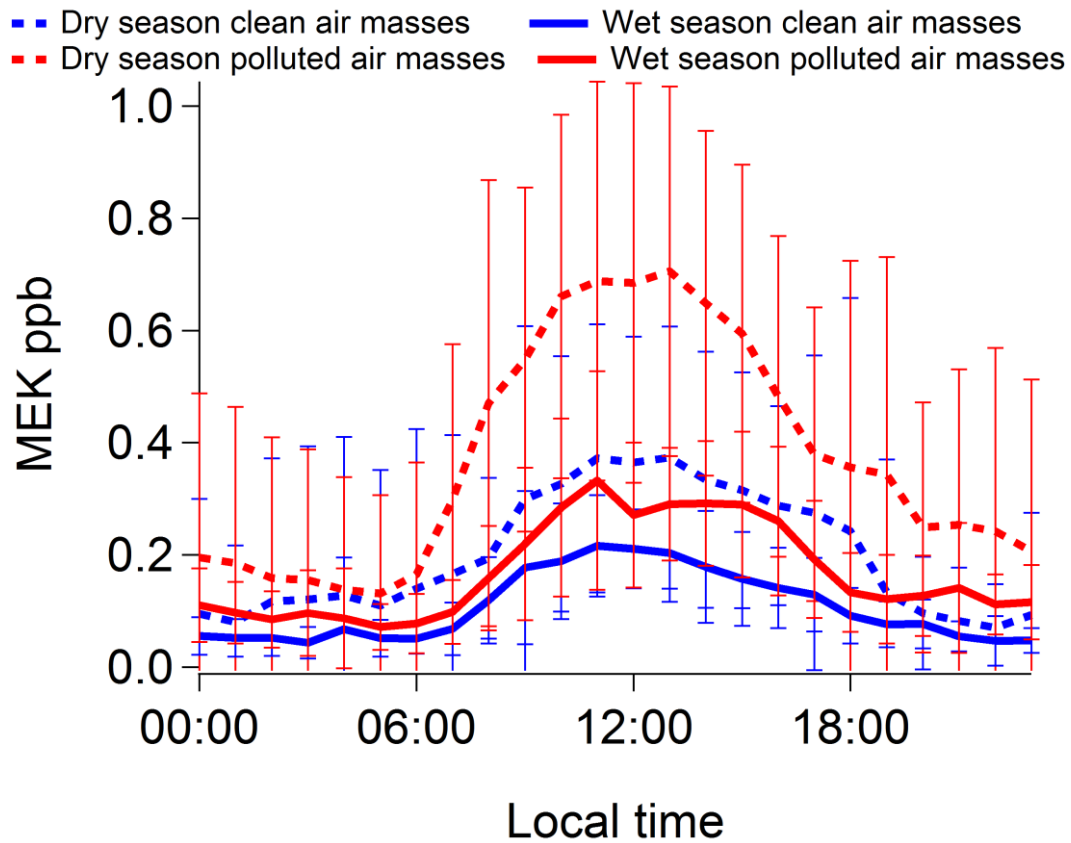


Figure 7: Hourly average concentrations of MEK in ppb for the clean conditions (blue) and the polluted conditions (red) at the T2 site. Dashed lines represent the dry season and thick lines represent the wet season. Error bars represent the standard deviation.

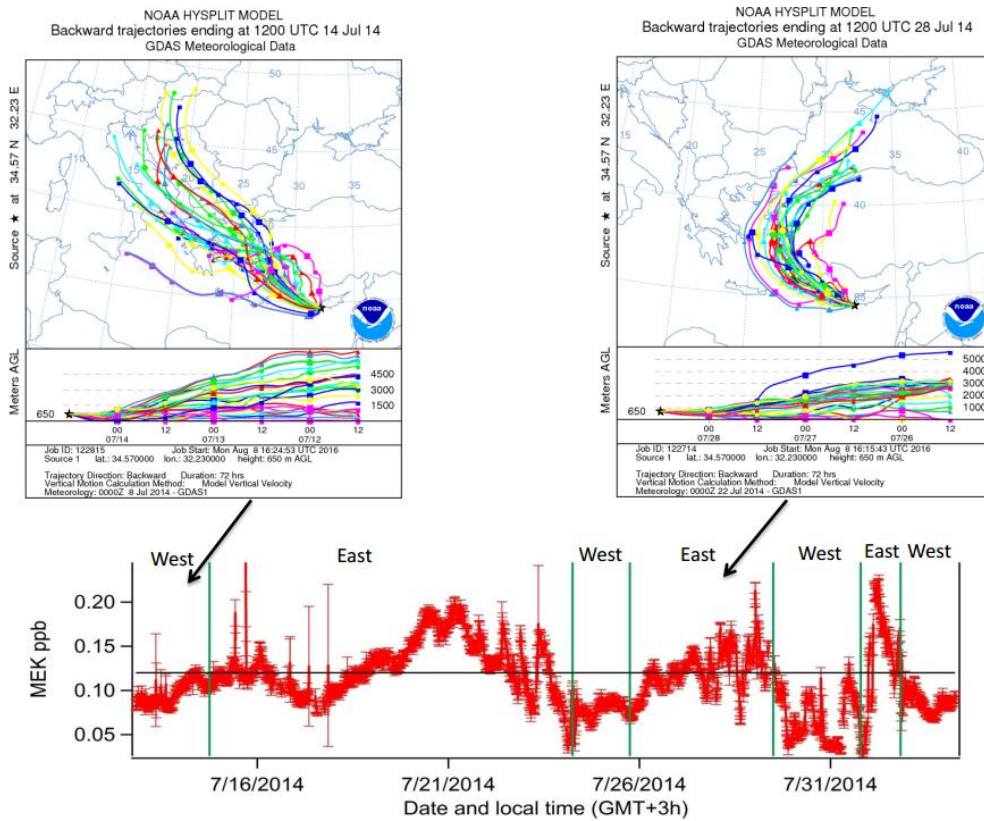


Figure 8: Timeline of MEK mixing ratios divided into periods when the air was coming from either Eastern or Western Europe. The HYSPLIT backward trajectories from 14 July and 28 August 2014 are shown based on the origin of the air masses. The black line represents the average of the whole campaign.