

1 Dear Dr. Robert Harley,

2 Thank you very much for your decision on our paper. We made the correction about how to deal with the data below
3 the limits of detection and addressed them as below. Meanwhile, data availability was updated in the revised manuscript
4 as “The data used in this study are available to all scientists agreeing to the AQABA protocol at:
5 <http://doi.org/10.5281/zenodo.3974228> (Wang et al., 2020).”

6 **“Please can you consider one further point relating to handling of data below the limit of detection (LOD):**
7 **rather than excluding those points (which are still valid measurements) as you have done, is there a way you can**
8 **use some small non-zero value such as LOD/2 or the square root of the LOD or ..., to avoid biasing the mean by**
9 **systematically excluding all of the lowest measured values.”**

10 Thanks for the comment and suggestion. We now included the data below the limit of detection using actual values
11 determined by the data processing method as suggested by the reviewer. The sentence was revised in the text (2.2.3)
12 as “Data below LOD were kept as determined for statistic analysis (Figure 2 and Table 1).” Table 1 and Figure 2 were
13 updated by including the data below the LOD. Following text were also revised accordingly (revised parts were marked
14 in red):

15 **Line 18:** acetaldehyde (0.16 ppb over the Arabian Sea... “Revised to “acetaldehyde (0.13 ppb over the Arabian Sea...”.

16 **Line 219-222:** “Acetaldehyde was measured at relatively low mixing ratios over the Arabian Sea (0.16 ± 0.12 ppb,
17 median: 0.13 ppb), which is comparable than the levels reported by the measurements done in northern-hemisphere
18 open ocean (see, Table 2). Over the Gulf of Aden, acetone and MEK had slightly higher mixing ratios than those over
19 the Arabian Sea.” Revised to “Acetaldehyde was measured at relatively low mixing ratios over the Arabian Sea (0.13
20 ± 0.12 ppb, median: 0.09 ppb), which is comparable than the levels reported by the measurements done in northern-
21 hemisphere open ocean (see, Table 2). Over the Gulf of Aden, **acetaldehyde**, acetone and MEK had slightly higher
22 mixing ratios than those over the Arabian Sea.”.

23 **Line 381-383:** “The mixing ratios of unsaturated carbonyls were generally low with values below 30 ppt over the
24 Mediterranean Sea and the clean 381 regions (the Arabian Sea and the Gulf of Aden, 12 - 21 ppt). The Red Sea region
25 and the Gulf of Oman had slightly higher levels 382 (13 – 60 ppt). The highest values were again observed in the
26 Arabian Gulf (25 – 115 ppt) followed by Suez (11 – 68 ppt).” Revised to “The mixing ratios of unsaturated carbonyls
27 were generally **~ 10 ppt or lower than the LOD** over the Mediterranean Sea and the clean regions (the Arabian Sea and
28 the Gulf of Aden). The Red Sea region and the Gulf of Oman had slightly higher levels (**LOD – 40 ppt**). The highest
29 values were again observed in the Arabian Gulf (**20 – 110 ppt**) followed by Suez (**LOD – 60 ppt**).”.

30 **Line 398-401:** “Around the Arabian Peninsula, the level of aromatic carbonyls declined with increasing carbon number
31 over most of the regions except in the Red Sea South where C8 carbonyls were slightly higher than C7 (Figure 2).
32 Interestingly, only in the Suez region, were the C7 aromatic carbonyls more abundant than other aromatic carbonyls,
33 whereby the mean value (128 ± 229 ppt) was much higher than the median value (30 ppt),” Revised to “Around the
34 Arabian Peninsula, the level of aromatic carbonyls declined with increasing carbon number over most of the regions
35 except in the Red Sea South, **Gulf of Oman and Arabian Gulf where C7 carbonyls were comparable to C8 carbonyls**
36 (Figure 2). Interestingly, only in the Suez region, were the C7 aromatic carbonyls more abundant than other aromatic
37 carbonyls, whereby the mean value (**90 ± 200 ppt**) was much higher than the median value (**20 ppt**).”

38 Best regards,

39 Nijing Wang

Measurements of carbonyl compounds around the Arabian Peninsula: overview and model comparison

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Abstract

Volatile organic compounds (VOCs) were measured around the Arabian Peninsula using a research vessel during the AQABA campaign (Air Quality and Climate Change in the Arabian Basin) from June to August 2017. In this study we examine carbonyl compounds, measured by a proton transfer reaction mass spectrometer (PTR-ToF-MS), and present both a regional concentration distribution and a budget assessment for these key atmospheric species. Among the aliphatic carbonyls, acetone had the highest mixing ratios in most of the regions traversed, varying from 0.43 ppb over the Arabian Sea to 4.5 ppb over the Arabian Gulf, followed by formaldehyde (measured by Hantzsch monitor, 0.82 ppb over the Arabian Sea and 3.8 ppb over the Arabian Gulf) and acetaldehyde (0.46-1.3 ppb over the Arabian Sea and 1.7 ppb over the Arabian Gulf). Unsaturated carbonyls (C4-C9) varied from 10 to 700 ppt during the campaign, and followed similar regional mixing ratio dependence as aliphatic carbonyls, which were identified as oxidation products of cycloalkanes over polluted areas. We compared the measurements of acetaldehyde, acetone and methyl ethyl ketone to global chemistry-transport model (EMAC) results. A significant discrepancy was found for acetaldehyde, with the model underestimating the measured acetaldehyde mixing ratio by up to an order of magnitude. Implementing a photolytically driven marine source of acetaldehyde significantly improved the agreement between measurements and model, particularly over the remote regions (e.g. Arabian Sea). However, the newly introduced acetaldehyde source was still insufficient to describe the observations over the most polluted regions (Arabian Gulf and Suez), where model underestimation of primary emissions and biomass burning events are possible reasons.

75 1 Introduction

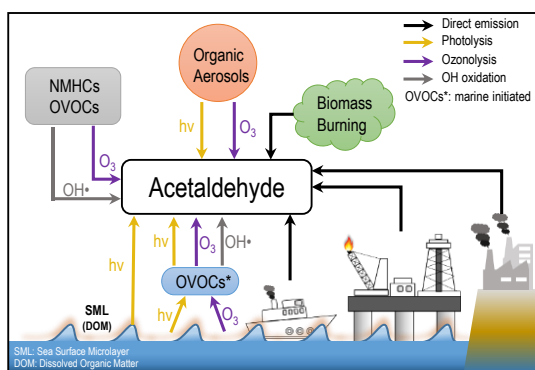
76 Carbonyl compounds (aldehydes and ketones) can be released into the air directly from a variety of primary biogenic and
77 anthropogenic sources. These include biomass burning (Holzinger et al., 1999; Holzinger et al., 2005; Koss et al., 2018), fossil fuel
78 combustion (Reda et al., 2014; Huang et al., 2018) including vehicles (Erickson et al., 2014; Dong et al., 2014), industrial solvent
79 use (Kim et al., 2008), and natural sources including plants and plankton (Zhou and Mopper, 1997; Warneke et al., 1999; Jacob et
80 al., 2002; Fall, 2003; Williams et al., 2004; Bourtsoukidis et al., 2014). However, secondary production via the atmospheric
81 oxidation of hydrocarbons is considered to be more important for many carbonyl compounds including acetone and acetaldehyde
82 (Jacob et al., 2002; Millet et al., 2010).

83 Carbonyls have several important roles in the atmosphere. They form as stable intermediates directly after hydrocarbon oxidation
84 by hydroxyl radicals, O_3 or NO_3 , when the peroxy radicals initially formed react with each other (permutation reactions) or with
85 NO. Their production is linked to tropospheric ozone formation (Carlier et al., 1986) and their loss, through oxidation and
86 photolysis, is an important source of hydroxyl and hydroperoxyl radicals (HO_x) in the dry upper troposphere (Colomb et al., 2006).
87 Carbonyls serve as precursors of peroxyacetyl nitrates (PANs) which are important atmospheric NO_x (NO and NO_2) reservoir
88 species (Finlayson-Pitts and Pitts, 1997; Edwards et al., 2014; Williams et al., 2000). Carbonyl compounds are also important for
89 the growth of atmospheric particles (Kroll et al. 2005) thereby indirectly impacting the Earth's radiative balance. The atmospheric
90 lifetimes of carbonyl compounds varies considerably, from less than one day for acetaldehyde (Millet et al., 2010) to more than 15
91 days for acetone (Jacob et al., 2002; Khan et al., 2015) in terms of tropospheric mean lifetime. A multiday lifetime means that
92 carbonyl compounds can impact the air chemistry on local, regional and even hemispheric scales. The numerous primary and
93 secondary sources of carbonyl compounds, as well as their multiple loss routes (photolysis, OH, NO_3 and O_3 oxidation) makes
94 budget assessments difficult.

95 The most predominant atmospheric carbonyl compounds besides formaldehyde are acetaldehyde and acetone. They have been
96 reported to vary from tens or hundreds of ppt in remote areas (Warneke and de Gouw, 2001; Wisthaler, 2002; Lewis et al.,
97 2005; White et al., 2008; Colomb et al., 2009; Read et al., 2012; Sjostedt et al., 2012; Tanimoto et al., 2014; Yang et al.,
98 2014; Hornbrook et al., 2016; Wang et al., 2019) to several ppb in urban and polluted areas (Dolgorouky et al., 2012; Guo et al.,
99 2013; Stoeckenius and McNally, 2014; Koss et al., 2015; Sahu et al., 2017; Sheng et al., 2018). Generally, secondary photochemical
100 formation from various precursors is the main source for those carbonyl compounds. However, several recent studies have shown
101 that acetaldehyde mixing ratios in both the remote marine boundary layer and the free troposphere could not be explained by
102 known photochemistry as implemented in various atmospheric chemistry models, which consistently underestimated the
103 measurements by an order of magnitude or more (Singh et al., 2003; Read et al., 2012; Wang et al., 2019). Several potential
104 additional acetaldehyde sources have been proposed including new hydrocarbon oxidation mechanisms, aerosol related sources
105 and oceanic sources. One possible source of acetaldehyde in the remote marine boundary layer is oceanic emission from the photo
106 degradation of colored dissolved organic matter (CDOM) in sea-surface water, where acetaldehyde could be produced together
107 with other low molecular weight carbonyl compounds (Kieber et al., 1990; Zhou and Mopper, 1997; Sinha et al., 2007; Dixon et al.,
108 2013). Nevertheless, due to both limited airborne and seawater measurements of acetaldehyde, the importance of oceanic emission
109 is still under debate (Millet et al., 2010; Wang et al., 2019). In order to better understand the atmospheric budgets of acetaldehyde
110 (and the other carbonyl compounds), it is informative to analyze a dataset of multiple carbonyl compounds in both polluted and
111 clean environments, with influence from marine emissions, varying particulate loadings, and high rates of oxidation as shown in
112 Figure 1, which demonstrates the main formation pathways of acetaldehyde during this campaign.

113 During the shipborne research campaign AQABA, carbonyl compounds were continuously measured by PTR-ToF-MS onboard a
 114 research vessel that circumnavigated the Arabian Peninsula. During the campaign, chemically distinct air masses were sampled,
 115 which had been influenced by primary emissions of hydrocarbons and inorganic pollutants (NO_x , SO_2 and CO) from petroleum
 116 industries and marine transport (Bourtsoukidis et al., 2019; Celik et al., 2019), by pollution from urban areas (Pfannerstill et al.,
 117 2019), and clean marine influenced air (Edtbauer et al., 2020). It is a unique dataset of carbonyl compounds encompassing starkly
 118 different environmental conditions from a region with few (or none) available in-situ measurements to date.

119 In this study, we provide an overview of carbonyl compound mixing ratios (aliphatic, unsaturated and aromatic) over the
 120 Mediterranean Sea, Suez, Red Sea, Arabian Sea and Arabian Gulf. Using an empirical method based on measured hydrocarbon
 121 precursors, we have analyzed the relative importance of the photochemical sources for the carbonyl compounds observed. The
 122 analysis is then extended to include sources and transport by using a global model EMAC (5th generation European Centre –
 123 Hamburg general model, ECHAM5 coupled to the modular earth submodel system, MESSy, applied to atmospheric chemistry).
 124 Model measurement differences are investigated in both clean and polluted regions, with particular emphasis on acetaldehyde.



125
 126 Figure 1. Diagram of possible sources and formation pathways of acetaldehyde during the AQABA campaign.
 127

128 2 Methods

129 2.1 AQABA campaign

130 The AQABA campaign was conducted onboard the research vessel Kommandor Iona (KI) from the end of June to the end of
 131 August 2017. The ship started from Southern France, proceeded across the Mediterranean, through the Suez Canal, around the
 132 Arabian Peninsula into the Arabian Gulf and on to Kuwait, thereafter returning along the same route. Five laboratory containers
 133 were loaded onto the vessel, containing multiple gas and particle phase measurement instruments as well as a weather station.

134 2.2 PTR-ToF-MS

135 2.2.1 Sampling and instrument set-up

136 A high-flow inlet (stainless steel tubing, 0.2 m diameter, 5.5 m tall and 3 m above the top of the containers and the front deck) was
 137 installed at the front of the ship where the laboratory containers were located. A high flow of air (approximately $10 \text{ m}^3 \text{ min}^{-1}$) was
 138 drawn through the inlet which provided a common attachment point for sub-sampling lines for all gas-phase measurement
 139 instruments. An air flow of 5 standard L min^{-1} for the first leg and 3.5 standard L min^{-1} for the second leg was pumped into the

140 on-board lab container through an 1/8" (O.D. = 1.27cm) FEP (fluorinated ethylene propylene) tubing (about 10 m long) insulated
141 and heated to 50-60 °C. A PTFE (polytetrafluoroethylene) filter was placed at the beginning of the inlet to prevent insects, dust
142 and particles entering the instruments. Every 2-5 days, the filter was replaced depending on the degree of pollution encountered.
143 Inside the VOC instrument container, the PTR-ToF-MS (8000, Ionicon Analytik GmbH Innsbruck, Austria) sampled a sub-flow
144 at 80-100 sccm through 1/8" (0.3175 cm) FEP tubing (~ 10 m in length, insulated and heated to 60 °C) from the main fast air flow
145 and then to the instrument's PEEK (polyether ether ketone) inlet which was likewise heated to 60 °C. The inlet system was shared
146 with total OH reactivity measurement (Pfannerstill et al., 2019).

147 The working principle of PTR-MS has been described in detail in previous studies (Lindinger et al., 1998; Ellis and Mayhew,
148 2013; Yuan et al., 2017). In brief, H₃O⁺ primary ions are generated in the ion source, and then drawn into the drift tube where they
149 interact with sampled ambient air. Inside the drift tube, VOCs with a proton affinity greater than that of H₂O (691 kJ mol⁻¹) are
150 protonated by proton transfer from H₃O⁺. The resulting secondary ions are transferred to the detector, in this case a time-of-flight
151 mass spectrometer with mass resolution around 3500 for the first leg and 4500 for the second leg at mass 96amu. An internal
152 standard of trichlorobenzene (C₆H₃Cl₃) was continuously introduced into the instrument to ensure accurate mass calibration. Every
153 minute a spectrum with mass range (m/z) 0-450 was generated. The data reported in this study are all at 1 minute resolution unless
154 otherwise specified.

155 2.2.2 Instrument characterization

156 The instrument background was determined every three hours for 10 minutes with synthetic air. 4-point calibrations were
157 performed five times during the whole campaign using a standard gas mixture (Apel-Riemer Environmental inc., Broomfield, USA)
158 containing 14 compounds (methanol, acetonitrile, acetaldehyde, acetone, dimethyl sulfide, isoprene, methyl vinyl ketone,
159 methacrolein, methyl ethyl ketone, benzene, toluene, xylene, 1,3,5-trimethylbenzene and α -pinene). It has been previously reported
160 that the sensitivity of some compounds measured by PTR-MS are humidity dependent (de Gouw and Warneke, 2007). As the
161 relative humidity (RH) was expected to be high and varying (marine boundary layer with occasional desert air influence), humidity
162 calibration was combined with 4-point calibration by humidifying the gas mixture at different levels from 0% - 100% RH.

163 2.2.3 Data analysis

164 The data were initially processed by the PTR Analyzer software (Müller et al., 2013) to identify and integrate the peaks. After
165 obtaining the raw data (counts per second for each mass identified), a custom-developed python-based program was used to further
166 process the data to final mixing ratios. For compounds present in the standard gas cylinder, interpolated sensitivities based on the
167 five in-campaign calibrations were applied to derive the mixing ratios; while mixing ratios of the other masses were calculated by
168 using a proton transfer reaction rate constant (k_{PTR}) of 2.0×10^{-9} cm³ s⁻¹. The uncertainty associated with the mixing ratios of the
169 calibrated compounds was around 6-17% (see Table S1). For the mixing ratios derived by assuming k_{PTR} , the accuracy was around
170 $\pm 50\%$ (Zhao and Zhang, 2004). The detection limit (LOD) was calculated from the background measurement with 3 times the
171 standard deviation (3σ), 52 ± 26 ppt for acetaldehyde, 22 ± 9 ppt for acetone and 9 ± 6 ppt for methyl ethyl ketone (MEK) (Table
172 S1). Data below LOD were ~~kept as determined for further statistic analysis (Figure 2 and Table 1) excluded from the data set instead~~
173 ~~of giving zero.~~

174 In this study, we have interpreted ion masses with the exact masses corresponding to C_nH_{2n}O, C_nH_{2n-2}O and C_nH_{2n-8}O as aliphatic,
175 unsaturated and aromatic carbonyls, respectively (see exact protonated m/z in Table S2). Carbonyl compounds with a carbon
176 number of three and above can be either aldehydes or ketones, which are not distinguishable with PTR-ToF-MS using H₃O⁺ as the

177 primary ion. However, laboratory experiments have shown that protonated aldehydic ions with carbon atoms more than three tend
178 to lose a H₂O molecule and fragment to other masses (Buhr et al., 2002;Spanel et al., 2002). Moreover, although both ketones and
179 aldehydes can be produced via atmospheric oxidation processes, ketones tend to have longer atmospheric lifetimes and higher
180 photochemical yields than aldehydes as mentioned in the introduction. The ratio of measured propanal to acetone was 0.07 in the
181 western Pacific coastal region (Schlundt et al., 2017), 0.06 in an urban Los Angeles (Borbon et al., 2013) and 0.17 - 0.22 in oil & gas
182 production regions (summarized by Koss et al., 2017). Therefore, signals on the exact mass of carbonyl compounds from the PTR-
183 ToF-MS are expected to be dominated by ketones, particularly in regions remote from the sources.

184 **2.3 Meteorological data and other trace gases**

185 The meteorological data were obtained by using a commercial weather-station (Sterela) which monitored wind speed, wind
186 direction, relative humidity (RH), temperature, speed of the vessel, and GPS etc. The actinic flux was measured by a spectral
187 radiometer (Metcon GmbH; Meusel et al., 2016). Non methane hydrocarbons (NMHC) mixing ratios were measured by a gas
188 chromatograph with flame ionization detector (GC-FID) online with the time resolution of 50 minutes. It measured hydrocarbons
189 (C₂-C₈) and aromatics (C₆-C₈) with the average LOD < 10 ppt for most of compounds. For a detailed instrumental description
190 see Bourtsoukidis et al. (2019). Formaldehyde mixing ratios were determined by a modified and optimized version of the
191 commercially available AL4021 (Aero-Laser, Germany), which utilizes the Hantzsch technique (Stickler et al., 2006). Methane
192 and carbon monoxide (CO) levels were monitored by a cavity ring-down spectroscopy analyzer (Picarro G2401). Ozone was
193 measured with an absorption photometer (Model 202 Ozone Monitor, 2B Technologies, Boulder, Colorado). Due to the potential
194 interference from sampling our own ship exhaust in which carbonyl compounds may be present (Reda et al., 2014), a filter was
195 applied to the data set based on the wind direction and NO_x, SO₂ and ethene levels.

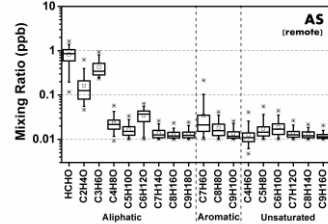
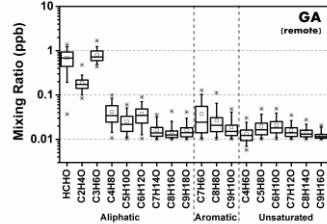
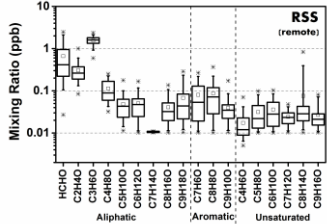
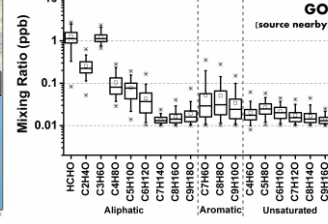
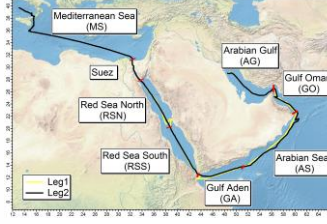
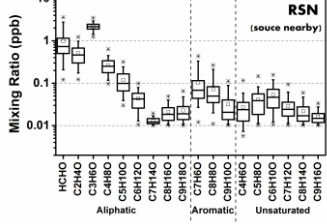
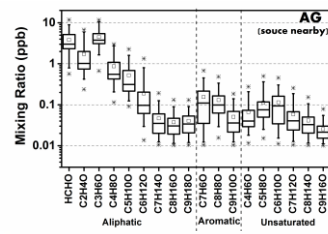
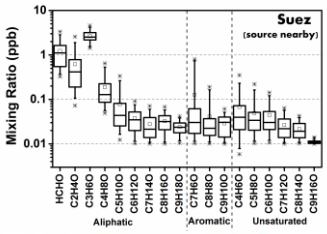
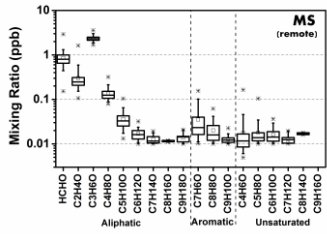
196 **2.4 Model simulations**

197 The EMAC (ECHAM5/MESy Atmospheric Chemistry) model was used to simulate atmospheric mixing ratios of several
198 carbonyl compounds along the cruise track covered during the AQABA campaign. The EMAC model is an atmospheric chemistry-
199 general circulation model simulating the process of tropospheric air by considering processes which could influence trace gases
200 mixing ratios, such as transport, chemistry, interaction with ocean/land, dry deposition and so on (Pozzer et al., 2007;Pozzer et al.,
201 2012;Lelieveld et al., 2016). The model applied in this study is a combination of the 5th generation of European Centre Hamburg
202 general circulation model (ECHAM5) (Roeckner et al., 2006) and the 2nd version of Modular Earth Submodel System (MESSy2)
203 (Jöckel et al., 2010), where a comprehensive chemistry mechanism MOM (Mainz Organic Mechanism) was deployed (Sander et
204 al., 2019). The model considers direct emissions (such as anthropogenic, biogenic, biomass burning etc.), atmospheric transport
205 and mixing, photochemical production of carbonyls (by OH, O₃ and NO₃), as well as physical and chemical removal processes.
206 The global fire assimilation system was used for biomass burning emissions (Kaiser et al., 2012). The exchange of organic
207 compounds between ocean and atmosphere was considered in EMAC via the AIRSEA submodel, described in detail in Pozzer et
208 al. (2006). The transfer velocity is calculated online and the concentration in the water is prescribed by the user. For acetone, a
209 constant water concentration of 15 nmol/L is used, following the suggestion of Fischer et al. (2012). The model configuration in
210 the study is the same as the model applied in Bourtsoukidis et al. (2020), where a natural non-methane hydrocarbon source (ethane
211 and propane) was implemented. The model is in the resolution of T106L31 (i.e. ~ 1.1° × 1.1° horizontal resolution and , 31 vertical
212 hybrid pressure levels up to 10 hPa) and the time resolution of 10 minutes. The measurement data of PTR-ToF-MS were averaged
213 to 10-minute resolution to match the model data resolution for further comparison.

214

215 3 Results and discussion

216 Around the Arabian Peninsula, the mixing ratios of individual carbonyl compounds varied over a wide range, from tens of ppt to
217 ppb levels. In this study, we divided the dataset geographically into eight regions (Figure 2, middle graph) to classify and
218 characterize the primary and secondary origins of carbonyl compounds. The regional delineations were: the Mediterranean Sea
219 (MS), Suez, Red Sea North (RSN), Red Sea South (RSS), Gulf of Aden (GA), Arabian Sea (AS), Gulf of Oman (GO) and Arabian
220 Gulf (AG), the same as those described by Bourtsoukidis et al. (2019). Figure 2 shows the abundance of aliphatic, aromatic and
221 unsaturated carbonyl compounds (carbonyls) for each region. Generally, aliphatic carbonyls were present at much higher mixing
222 ratios than aromatic and unsaturated carbonyls, with smaller carbonyl compounds (formaldehyde, acetaldehyde, C3 and C4
223 carbonyls) dominating the distribution. The mixing ratios of aliphatic carbonyls decreased dramatically from C5 carbonyls with
224 increasing carbon number. The box plots (Figure 2) also show that carbonyl compounds were measured at higher mixing ratios
225 and were more variable over Suez region and the Arabian Gulf. The abundance of carbonyl compounds varied markedly from
226 region to region with highest and lowest values found in the Arabian Gulf and the Arabian Sea, respectively. Table 1 shows the
227 mean, standard deviation and the median values for carbonyls in each region. In the following sections, each class of carbonyl
228 compounds are investigated in greater detail.



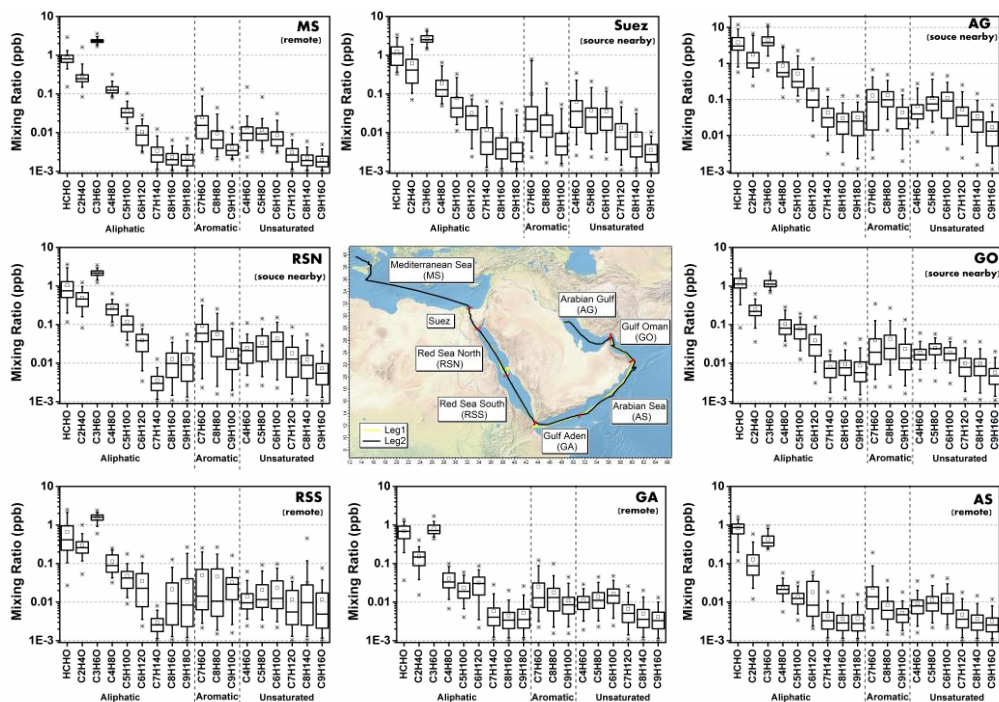


Figure 2. Overview of mixing ratios for aliphatic, aromatic and unsaturated carbonyl compounds (CxHyO). The boxes represent 25% to 75% of the data with the central line and square indicating the median and the mean values, respectively. The whiskers show data from 5% to 95% and stars were drawn for the minimum and maximum data points within 1% to 99% of the dataset. Within brackets under the region acronyms the main characteristics of the air masses are indicated, based on non-methane hydrocarbon variability-lifetime results (b factor) from Bourtsoukidis et al. (2019). The data used for map plotting was from public domain GIS data found on the Natural Earth web site (<http://www.naturalearthdata.com>) and was read into Igor using the IgorGIS XOP beta.

3.1 Aliphatic carbonyls (C_nH_{2n}O)

3.1.1 Overview

Relatively high mean mixing ratios of aliphatic carbonyls were observed over the Arabian Gulf; the highest being acetone (C3 carbonyl compound) at 4.50 ± 2.40 ppb (median: 3.77 ppb), followed by formaldehyde at 3.83 ± 2.55 ppb (median: 3.02 ppb), acetaldehyde at 1.73 ± 1.61 ppb (median: 1.02 ppb) and MEK (C4 carbonyl compound) at 0.87 ± 0.71 ppb (median: 0.56 ppb). As the Arabian Gulf is highly impacted by the oil and gas industry, we compared the measurements of the four aforementioned carbonyl compounds with those measured in the oil and gas region (Table 2). Except for formaldehyde, acetaldehyde, acetone and MEK were lower than the mixing ratios measured in the Uintah Basin, which was influenced by intensive oil and natural gas activities (Koss et al., 2015). The general distribution of the aliphatic carbonyls in the Uintah Basin is similar to the Arabian Gulf, with acetone levels being approximately twice as those of acetaldehyde. The carbonyl mixing ratios in the Arabian Gulf were comparable to those measured in Hickory (PA, USA) surrounded by natural gas wells (Swarthout et al., 2015). Koss et al. (2017) reported the max boundary layer enhancement of carbonyl compounds (C2-C7) measured during an aircraft measurement above the most productive oil field in the United States (Permian Basin). Within the boundary layer of the Permian Basin, C5-C7 aliphatic

251 carbonyls had mixing ratios of 0.34 ppb, 0.08 ppb and 0.03 ppb; which are of the same magnitude but lower than the levels
252 measured over the Arabian Gulf for C5 (0.52 ± 0.48 ppb), C6 (0.19 ± 0.25 ppb) and C7 ($0.05-04 \pm 0.04$ ppb) carbonyl compounds.
253 The sources of the major carbonyls in the Arabian Gulf will be discussed in details in section 3.1.2 and 3.4.3.

254 In contrast, aliphatic carbonyls had much lower average mixing ratios over the Arabian Sea and the Gulf of Aden especially for
255 C7-C9 carbonyls with mean mixing ratios below the detection limit for most of the time. During the summertime AQABA
256 campaign, the prevailing wind direction over the Arabian Sea was southwest (Figure S1). Four-day back trajectories indicate the
257 air was transported from the Arabian Sea (Northwestern Indian Ocean), passing East Africa coast, which brought relatively clean,
258 photochemically aged airmasses (Bourtsoukidis et al., 2019). The mean level of acetone over the Arabian Sea (0.43 ± 0.18 ppb,
259 median: 0.34 ppb) is close to the level measured in the marine boundary layer of Western Indian Ocean (0.49 ppb) (Warneke and
260 de Gouw, 2001) and comparable to other reported values from open-sea air measurement (see Table 2). Acetaldehyde was
261 measured at relatively low mixing ratios over the Arabian Sea ($0.46-13 \pm 0.12$ ppb, median: $0.43-09$ ppb), which is comparable
262 than the levels reported by the measurements done in northern-hemisphere open ocean (see, Table 2). Over the Gulf of Aden,
263 acetaldehyde, acetone and MEK had slightly higher mixing ratios than those over the Arabian Sea.

264 The Mediterranean Sea had somewhat higher levels of aliphatic carbonyls than the clean regions (the Arabian Sea and the Gulf of
265 Aden) but with acetone (above 2ppb) dominating the distribution. Much higher acetone level than acetaldehyde level was also
266 observed for some costal site measurement which was impacted by continental air (White et al., 2008; Schlundt et al., 2017, see
267 Table 2). Larger aliphatic carbonyls (C6-C9) were below the detection limit most of the time. The aliphatic carbonyls levels over
268 the Gulf of Oman were higher than the clean regions, while C1-C5 carbonyls were more variable over the Gulf of Oman compared
269 to those over the Mediterranean Sea. This is probably because the Gulf of Oman connects to the Arabian Gulf where intense oil
270 and gas industrial activities are located. Over the Gulf of Oman, polluted air from the nearby sources of the Arabian Gulf is
271 occasionally mixed with the clean air from the open sea (the Arabian Sea) under southeast wind conditions (Figure S1).

272 Another region where abundant aliphatic carbonyls were observed was Suez region. The air in this region was mainly influenced
273 by nearby cities and marine transportation (ship emissions within the Suez Chanel) (Bourtsoukidis et al., 2019;Pfanterstill et al.,
274 2019). Therefore abundant precursors were available in Suez region, producing more carbonyls regionally especially for shorter-
275 lived compounds (formaldehyde and acetaldehyde). Besides the local-scale emissions and photochemical production contribution
276 to the carbonyls over Suez, the longer lived carbonyls (e.g. acetone) could be also transported from the Mediterranean Sea (where
277 acetone was high). Four-day back trajectories indicate the air reaching Suez region was mostly originated from Europe continent
278 passing over the Mediterranean Sea (Bourtsoukidis et al., 2019). Meanwhile, ocean uptake of acetone from the air due to polluted
279 continental outflow (Marandino et al., 2005) as well as dilution and mixing with free tropospheric air during transport can modulate
280 acetone mixing ratios. Although the mean mixing ratios of aliphatic carbonyls over Suez were much lower than those over the
281 Arabian Gulf, the variations were still more significant than other regions (not including the Arabian Gulf, see Table 1).

282 Over the Red Sea, acetone was the most abundant aliphatic carbonyls followed by formaldehyde and acetaldehyde. The mixing
283 ratios of acetaldehyde and acetone over the northern part of the Red Sea were similar to those levels measured in western Pacific
284 coastal regions (South China Sea, Table 2). It is worth noticing that the levels of aliphatic carbonyls in the northern part of the Red
285 Sea were almost two times higher than the southern part of the Red Sea. According to the four-day back trajectories reported by
286 Bourtsoukidis et al. (2019), the measured air masses travelled to the northern part was from southern Europe and northeast Africa
287 while the southern part was more influenced by air from the northern part of the Red Sea mixed with the air masses from desertic
288 areas of central Africa. Therefore, less primary precursors as well as carbonyls were transported to the southern part of the Red

289 Sea compared to the northern part. Moreover, the unexpected sources of hydrocarbons (ethane and propane) from Northern Red
 290 Sea deep water reported by Boursoukidis et al. (2020) would lead to higher carbonyl levels in the Northern part compared with the
 291 Southern part due to the additional precursors in the Red Sea North. However, acetaldehyde was still found to be significantly
 292 underestimated compared to the model results, even taking the deep-water source into consideration (section 3.3). This indicates
 293 that extra sources of acetaldehyde may exist, which will be discussed in detail in section 3.4.

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Table 1. Mean, standard deviation (SD) and median mixing ratios of aliphatic, unsaturated and aromatic carbonyls in different regions.

		Aliphatic Carbonyls								
		HCHO	CH3CHO	C3H6O	C4H8O	C5H10O	C6H12O	C7H14O	C8H16O	C9H18O
MS	mean	0.86	0.30	2.37	0.14	0.04	0.02	0.01	0.01	0.01
	SD	0.41	0.25	0.37	0.05	0.02	0.01	0.00	0.00	0.00
	median	0.80	0.25	2.32	0.12	0.03	0.02	0.01	0.01	0.01
Suez	mean	1.23	0.62	2.64	0.19	0.08	0.04	0.03	0.03	0.02
	SD	0.76	0.58	0.77	0.15	0.08	0.02	0.02	0.02	0.01
	median	1.11	0.42	2.52	0.13	0.04	0.04	0.02	0.03	0.02
RSN	mean	0.99	0.51	2.17	0.27	0.12	0.04	0.01	0.02	0.02
	SD	0.78	0.26	0.45	0.11	0.07	0.02	0.00	0.01	0.01
	median	0.73	0.46	2.17	0.25	0.10	0.04	0.01	0.02	0.02
RSS	mean	0.66	0.31	1.56	0.11	0.05	0.05	0.01	0.04	0.07
	SD	0.62	0.17	0.38	0.06	0.03	0.03	0.00	0.03	0.07
	median	0.40	0.26	1.60	0.09	0.04	0.05	0.01	0.03	0.04
GA	mean	0.69	0.19	0.81	0.04	0.03	0.04	0.02	0.02	0.02

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	<u>mean</u>	0.09	0.03	< LOD	0.06	0.04	0.03	0.01	0.01	< LOD
<u>Suez</u>	<u>SD</u>	0.20	0.04	n.a.	0.08	0.04	0.03	0.01	0.01	n.a.
	<u>median</u>	0.02	0.01	< LOD	0.04	0.02	0.02	0.01	< LOD	< LOD
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	<u>mean</u>	0.09	0.05	0.02	0.03	0.03	0.04	0.02	0.01	0.01
<u>RSN</u>	<u>SD</u>	0.10	0.06	0.02	0.02	0.03	0.04	0.02	0.01	0.01
	<u>median</u>	0.06	0.04	0.01	0.02	0.03	0.03	0.01	0.01	< LOD
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	<u>mean</u>	0.05	0.04	0.03	0.01	0.02	0.02	0.01	0.03	0.01
<u>RSS</u>	<u>SD</u>	0.06	0.06	0.03	0.01	0.02	0.02	0.01	0.07	0.01
	<u>median</u>	0.01	0.01	0.03	0.01	0.01	0.01	< LOD	< LOD	< LOD
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	<u>mean</u>	0.02	0.02	0.01	0.01	0.01	0.02	0.01	< LOD	< LOD
<u>GA</u>	<u>SD</u>	0.03	0.02	0.01	0.01	0.01	0.01	0.01	n.a.	n.a.
	<u>median</u>	0.01	0.01	0.01	0.01	0.01	0.01	< LOD	< LOD	< LOD
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	<u>mean</u>	0.02	0.01	< LOD	0.01	0.01	0.01	< LOD	< LOD	< LOD
<u>AS</u>	<u>SD</u>	0.03	0.01	n.a.	0.01	0.01	0.01	n.a.	n.a.	n.a.
	<u>median</u>	0.01	0.01	< LOD	0.01	0.01	0.01	< LOD	< LOD	< LOD
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	<u>mean</u>	0.04	0.04	0.02	0.02	0.02	0.02	0.01	0.01	0.01
<u>GO</u>	<u>SD</u>	0.06	0.05	0.03	0.01	0.01	0.01	0.01	0.01	n.a.
	<u>median</u>	0.02	0.02	0.01	0.02	0.02	0.02	0.01	0.01	< LOD
<hr/>										
	<u>mean</u>	0.12	0.13	0.04	0.07	0.11	0.12	0.05	0.03	0.02
<u>AG</u>	<u>SD</u>	0.14	0.10	0.04	0.06	0.10	0.10	0.05	0.03	0.02
	<u>median</u>	0.08	0.10	0.03	0.04	0.07	0.09	0.04	0.03	0.01

< LOD: the mixing ratios were lower than the limit of detection.

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Table 2. Mixing ratios (ppb) of OVOCs reported in previous observation in literature

Locations	Lon./Lat.	Height	Time	Technique	Formaldehyde	Acetaldehyde	Acetone	MEK	Literature
Open sea		m							
Tropical Atlantic Ocean	10° N-0° N 35° W-5° E	18	Oct.-Nov.	PTR-MS	n.r.	n.r.	0.53	n.r.	(Williams et al., 2004)
Atlantic Ocean	50° N-50° S 10-60° W	18	Oct.-Nov.	PTR-MS	n.r.	0.18 (Northern H) 0.08 (Southern H)	0.6 (North) 0.2 (South)	n.r.	(Yang et al., 2014)
Western North Pacific Ocean	15-20° N 137° E	6.5-14	May	PTR-MS	n.r.	n.r.	0.20-0.70	n.r.	(Tanimoto et al., 2014)
Western Indian Ocean	12° N-5° S 43-55° E	15	Feb.-Mar.	PTR-MS	n.r.	n.r.	0.49	n.r.	(Warneke and de Gouw, 2001)
Indian Ocean	19° N-13° S 67-75° E	10	Mar.	PTR-MS	n.r.	0.32-0.42 (continental outflow) 0.18-0.21 (equatorial marine)	1.11-2.08 (continental outflow) 0.51-0.62 (equatorial marine)	n.r.	(Wisthaler, 2002)
Southern Indian Ocean	30° S-49° S 30-100° E	15	Dec.	PTR-MS	n.r.	0.12-0.52	0.42-1.08	n.r.	(Colomb et al., 2009)
Coastal									
Caribbean Sea	10-30° N 60-80° W	10	Oct.	HPLC	0.61	0.57	0.40	0.03	(Zhou and Mopper, 1993)
Cape Verde Atmospheric Observatory	16.86° N 24.87° W	10	2006 - 2011	GC-FID	n.r.	0.43 (0.19-0.67)	0.55 (0.23-0.91)	n.r.	(Read et al., 2012)
Appledore Island, USA	42.97° N 70.62° W	5	Jul.-Aug.	PTR-MS	n.r.	0.40	1.5	0.20	(White et al., 2008)
Mace Head, Ireland	53.3° N 9.9° W	25	Jul.-Sep.	GC-FID	n.r.	0.44 (0.12-2.12)	0.50 (0.16-1.67)	n.r.	(Lewis et al., 2005)
Canadian Archipelago	68-75° N 60-100° W	Ship cruise	Aug.-Sep.	PTR-MS	n.r.	n.r.	0.34	n.r.	(Sjostedt et al., 2012)
Barrow Arctic	71.30° N 156.77° W	6	Mar.-Apr.	TOGA		0.10 ± 0.20	0.90 ± 0.30	0.19 ± 0.05	(Hornbrook et al., 2016)
South China Sea, Sulu Sea	2° N-15° N 108-124° E	10	Nov.	GC-MS	n.r.	0.86	2.1	0.06	(Schlundt et al., 2017)
Oil & Gas									
Horse Pool site, Uintah Basin, USA		Ground site	2012 - 2013	PTR-MS	3.71	4.27	7.97	2.81	(Koss et al., 2015)
Central United State		<600	Mar. - April	ToF-CIMS	1.13 ^a	0.5	1.5	0.2	(Koss et al., 2017)
Eagle Mountain Lake site, Texas, USA		Ground site	June	PTR-MS	n.r.	n.r.	3.2 (1.2-6.7)	0.3 (0.09-0.85)	(Rutter et al., 2015)

Hickory, Pennsylvania, a, USA	Ground site	June	PTR-MS	n.r.	1.29 (0.28-2.03)	3.22 (1.45-4.99)	0.73 (0.4-0.97)	(Swarthout et al., 2015)
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n.r.: not reported in the literature.

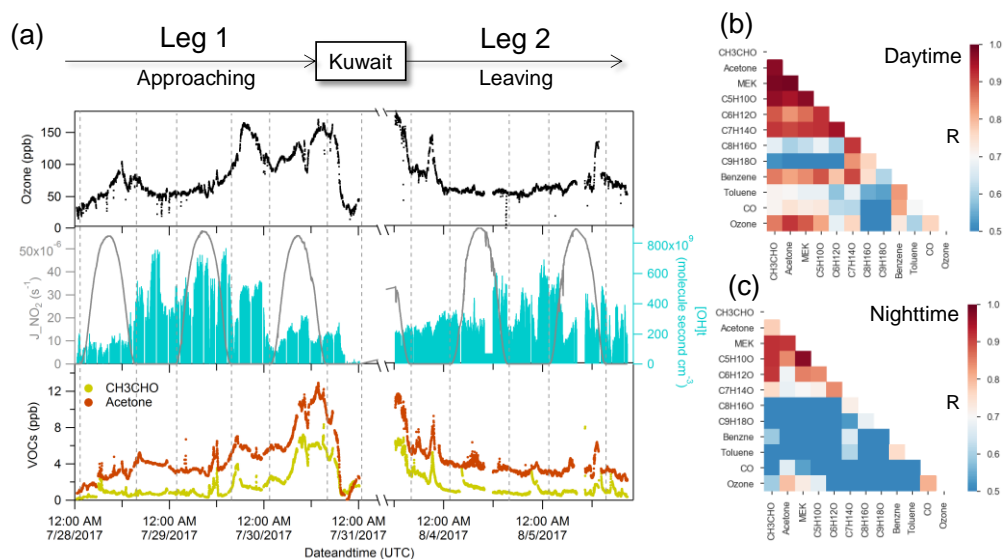
a: formaldehyde was measured by laser-induced fluorescence (LIF)

3.1.2 Case studies of polluted regions: the Arabian Gulf and Suez

The primary emission sources in the Arabian Gulf and Suez regions are quite different. While the Arabian Gulf is dominated by oil and gas operations, Suez is more influenced by ship emissions and urban areas (Bourtsoukidis et al., 2019). Carbonyl compounds were most abundant in these two areas. For further insight, we focused on a time series of selected trace-gases and their inter-correlations to better identify the sources of the major aliphatic carbonyls. Meanwhile, we calculated the OH exposure ($[OH]\Delta t$) based on hydrocarbon ratios (Roberts et al., 1984; de Gouw et al., 2005; Yuan et al., 2012) for the polluted regions (Arabian Gulf and Suez) where primary emissions have been identified (Bourtsoukidis et al., 2019; Bourtsoukidis et al. 2020), to better understand the photochemical aging of the major carbonyls using the following equation:

$$[OH]\Delta t = \frac{1}{k_X - k_Y} \cdot \left(\ln \frac{[X]}{[Y]} \Big|_{t=0} - \ln \frac{[X]}{[Y]} \right), \quad \text{Eq. (1)}$$

where X and Y refer to two hydrocarbon compounds with different rates of reaction with the OH radical (k). For this study, we chose toluene ($k_{OH+toluene}: 5.63E-12 \text{ cm}^3 \text{ molecule}^{-1}\text{s}^{-1}$) and benzene ($k_{OH+benzene}: 1.22E-12 \text{ cm}^3 \text{ molecule}^{-1}\text{s}^{-1}$) (Atkinson and Arey, 2003), because both compounds were measured by PTR-ToF-MS at high frequency and these values showed a good agreement with values measured by GC-FID (Figure S2). The approach detailed by Yuan et al. (2012) was applied to determine the initial emission ratio $\frac{[X]}{[Y]} \Big|_{t=0}$ in those two regions by only including nighttime data of benzene and toluene. We obtained initial emission ratios (toluene to benzene ratio) of 1.38 for the Arabian Gulf and 2.12 for the Suez region. Koss et al. (2017) summarized the toluene to benzene ratios observed in various locations and showed that urban and vehicle sources tend to have higher toluene to benzene ratio (mean ~ 2.5) than the ratios of oil & gas sources (mean ~ 1.2). Therefore, the toluene to benzene ratios obtained for those two regions agreed well with other studies done with similar emissions sources. The corresponding correlation plots of toluene and benzene for those two regions can be found in Figure S3.



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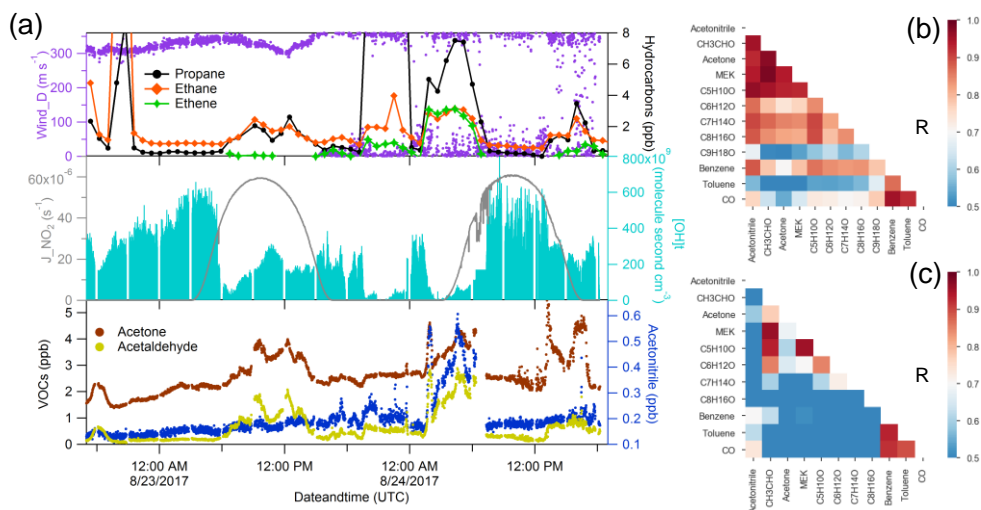
375 Figure 3. Case study of the Arabian Gulf. (a) Time series of selected species measured over the Arabian Gulf; (b) daytime
 376 correlation heat map of selected species; (c) nighttime correlation heat map of selected species.

377

378 Figure 3(a) shows the time series of acetaldehyde and acetone over the Arabian Gulf along with OH exposure ($[OH]t$) and ozone.
 379 We further separated the data into daytime and nighttime and calculated correlations among the carbonyls and other selected
 380 species (see Fig. 4b and c). Aliphatic carbonyls were well correlated with each other during the daytime and ozone had a generally
 381 good correlation with C2-C7 carbonyls ($r > 0.7$) during the daytime but a much lower correlation during the night, indicating ozone
 382 and carbonyls were co-produced via photochemical oxidation. Tadic et al. (2020) reported the net ozone production rate over the
 383 Arabian Gulf (32 ppb d^{-1}) was the greatest over the Arabian Peninsula. They show that strong ozone forming photochemistry
 384 occurred in this region, which would lead to abundant secondary photo-chemically produced products (including carbonyls).
 385 However, it should be noted the good correlation between ozone and carbonyls could in part be due to carbonyls co-emitted with
 386 ozone precursors (hydrocarbons) as primary emissions. In Figure 3 (a), the calculated OH exposure was high during the first night
 387 in leg 1, where an elevation of acetone mixing ratio was observed while the mixing ratio of acetaldehyde remained relatively
 388 constant. With limited OH radical abundance during the nighttime, the increased OH exposure indicates that the air reaching the
 389 ship was photochemically processed (aged). Therefore, the increase of acetone was mainly from long-distance transport as acetone
 390 has a much longer atmospheric lifetime than acetaldehyde. As the ship approached Kuwait, the calculated OH exposure was low
 391 (starting from 7/30/2017, 12:00 am UTC), which is an indicator of nearby emission sources. The lifetime of the OH radical derived
 392 from the measured OH reactivity also decreased from $\sim 0.1 \text{ s}$ to $\sim 0.04 \text{ s}$ during the same period (Pfannerstill et al., 2019). Oil
 393 fields and associated refineries are densely distributed in the northwest of the Arabian Gulf region (United States Central
 394 Intelligence Agency). The air reaching the ship when mixing ratios of acetone and acetaldehyde were highest was mainly from the
 395 Northwest (Iraq oil field region) according to the back trajectories (Bourtsoukidis et al., 2019). This suggests that the air masses
 396 encountered in Northwest Arabian Gulf were a combination of fresh emissions from nearby sources and photochemically processed

397 air transported from elsewhere. During the second leg, relatively low mixing ratios were identified in the same region (Northwest
 398 Arabian Gulf), which was mainly due to a greater influence of air masses originating from less populated desert regions of
 399 Northeast Iran (Bourtsoukidis et al., 2019) with much less influence from the oil field emissions, meaning less precursors were
 400 available for carbonyl production. Several plumes (extending over 2-3 hours) of elevated carbonyls with increased ozone were
 401 observed during the nighttime for both legs (Fig. 4a), indicating transport of highly polluted air.

402



403

404 Figure 4. Case study of Suez. (a) Time series of selected species measured over Suez; (b) correlation heat map of selected species
 405 during biomass burning plume (UTC 01:00 -06:00 August 24th 2017); (c) correlation heat map of selected species without the
 406 period of biomass burning plume.

407

408 For the Suez region (Gulf of Suez and Suez Canal), data were only available for the second leg. A significant increase of acetonitrile
 409 (over 400 ppt) was observed just before entering the Great Bitter Lake (see Figure 4a), indicating an increasing influence of biomass
 410 burning on the air composition (Lobert et al., 1990). Carbonyl compounds are important primary emissions in fresh biomass
 411 burning plumes (Holzinger et al., 1999;Schauer et al., 2001;Holzinger et al., 2001;Koss et al., 2018) as well as being formed as
 412 secondary products in more aged plumes (Holzinger et al., 2005). We further investigated the correlation coefficient among
 413 carbonyls during the biomass burning plume (Figure 4b) in Suez. Carbonyls had a high correlation with acetonitrile, benzene and
 414 among themselves, particularly for smaller carbonyls (acetaldehyde, C3-C5 carbonyls). The biomass burning emissions were
 415 probably transported by on the prevailing northerly wind (Figure S1) above Northeast Egypt where crop residues especially rice
 416 straw is often directly burned in the open fields (Abdelhady et al., 2014;Said et al., 2013;Youssef et al., 2009). Besides the direct
 417 biomass burning emission, the high mixing ratios and the good correlations of carbonyls could also have resulted from other
 418 sources as hydrocarbons (alkanes, alkenes and aromatics) which were elevated at the same time. Similar to conditions identified
 419 over the Arabian Gulf, elevated OH exposure accompanied with increasing acetone mixing ratio was observed during the first
 420 night over the Gulf of Suez, indicating aged air mass transportation. The OH exposure was then significantly lower during the
 421 daytime, when mixing ratios of carbonyls and alkanes increased as well. This indicates the presence of emission sources nearby.

422 Oil refineries located in the coastal side of Suez and oil tank terminals located in the northern part of the Gulf of Suez are likely
423 sources.

424 3.2 Unsaturated and aromatic carbonyls (C_nH_{2n-2}O), (C_nH_{2n-8}O)

425 3.2.1 Overview

426 The mixing ratios of unsaturated carbonyls were generally ~~low with values below ~ 30-10 ppt or lower than the LOD~~ over the
427 Mediterranean Sea and the clean regions (the Arabian Sea and the Gulf of Aden, ~~12-21 ppt~~). The Red Sea region and the Gulf of
428 Oman had slightly higher levels (~~13-LOD - 60-40 ppt~~). The highest values were again observed in the Arabian Gulf (~~25-20 - 115~~
429 ~~110 ppt~~) followed by Suez (~~11-LOD - 68-60 ppt~~). The numbers represent the range of the mean mixing ratios of unsaturated
430 carbonyls in each region. In terms of the mixing ratio distribution (Figure 2), the peak value was usually observed at C5 or C6
431 unsaturated carbonyls over most regions except for Suez where C4 carbonyl had the highest mixing ratio. Based on chemical
432 formulas, unsaturated carbonyls can be either cyclic carbonyl compounds or carbonyls containing a carbon-carbon double bond.
433 Therefore, the air chemistry could differ considerably depending on the compound assignment. A detailed analysis of the chemistry
434 of the unsaturated carbonyls measured will be given in the following section 3.2.2.

435 Regional variability was also observed for aromatic carbonyls with highest levels observed over the Arabian Gulf and Suez, and
436 much lower mixing ratios over the Arabian Sea, Mediterranean Sea and Gulf of Aden (Table 1). Several studies using PTR-MS
437 have reported values for m/z 107.049 (C7 aromatic carbonyls) attributed to benzaldehyde (Brilli et al., 2014; Koss et al., 2017; Koss
438 et al., 2018), m/z 121.065 (C8 aromatic carbonyls) attributed to toluene (Koss et al., 2018) or acetophenone (Brilli et al.,
439 2014) and m/z 135.080 (C9 aromatic carbonyls) attributed to methyl acetophenone (Koss et al., 2018) or benzyl methyl ketone
440 (Brilli et al., 2014) or 3,5-dimethylbenzaldehyde (Müller et al., 2012). Atmospheric aromatic carbonyls are produced via
441 photochemical oxidation of aromatic hydrocarbons (Finlayson-Pitts and Pitts Jr, 1999; Wyche et al., 2009; Müller et al., 2012) and
442 benzaldehyde was reported as having primary sources from biomass burning and anthropogenic emissions (Cabrera-Perez et al.,
443 2016). Around the Arabian Peninsula, the level of aromatic carbonyls declined with increasing carbon number over most of the
444 regions except in the Red Sea South ~~Gulf of Oman and Arabian Gulf~~ where ~~C8-C7 carbonyls were slightly higher than comparable~~
445 ~~to C7-C8 carbonyls~~ (Figure 2). Interestingly, only in the Suez region, were the C7 aromatic carbonyls ~~more abundant than other~~
446 aromatic carbonyls, whereby the mean value (~~128-90 ± 229-200 ppt~~) was much higher than the median value (~~30-20 ppt~~), indicating
447 strong primary sources of benzaldehyde in Suez. Otherwise, toluene was found to be more abundant over Suez with mean mixing
448 ratios of 271 ± 459 ppt than over other regions (the mean over the Arabian Gulf: 130 ± 160 ppt) which would also lead to higher
449 benzaldehyde as it is one of the OH-induced oxidation products of toluene via H-abstraction (Ji et al., 2017).

450 3.2.2 Potential precursors and sources of unsaturated carbonyls

451 Unsaturated carbonyls measured by PTR-MS have been only rarely reported in the atmosphere with the exception of methyl vinyl
452 ketone and methacrolein (C4 carbonyls) which are frequently reported as the oxidation products of isoprene (Williams et al., 2001;
453 Fan and Zhang, 2004; Wennberg et al., 2018). According to the GC-FID measurement, isoprene was below the detection limit for
454 most of the time during the AQABA cruise with the highest values observed in Suez (10 - 350 ppt). This shows that the AQABA
455 campaign was little influenced by either terrestrial or marine isoprene emissions. However, we observed unexpected high levels
456 on mass 69.070, which is usually interpreted as isoprene for PTR-MS measurements. Significant enhancements were even
457 identified while sampling our own ship exhaust (in PTR-MS but not GC-FID), suggesting the presence of an anthropogenic
458 interference at that mass under these extremely polluted conditions. Several studies have reported possible fragmentations of cyclic

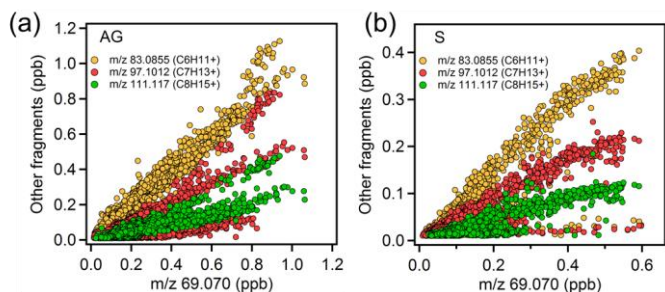
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459 alkanes giving mass (m/z) 69.070. These include: a laboratory study on gasoline hydrocarbon measurements by PTR-MS
 460 (Gueneron et al., 2015), a GC-PTR-MS study of an oil spill site combined with analysis of crude oil samples (Yuan et al., 2014)
 461 and an inter-comparison of PTR-MS and GC in an O&G industrial site (Warneke et al., 2014). From those studies, other
 462 fragmentations from C5-C9 cycloalkanes including m/z 43, m/z 57, m/z 83, m/z 111 and m/z 125 were identified together with
 463 m/z 69. Cyclic alkanes were directly measured in oil and gas fields (Simpson et al., 2010; Gilman et al., 2013; Li et al., 2017; Aklilu
 464 et al., 2018), vehicle exhaust (Gentner et al., 2012; Erickson et al., 2014), vessel exhaust (Xiao et al., 2018), accounting for a non-
 465 negligible amount of the total VOCs mass depending on the fuel type. Koss et al. (2017) reported enhancement of cyclic alkane
 466 fragment signals and increased levels of unsaturated carbonyls measured by PTR-ToF-MS over O&G region in the US. The
 467 unsaturated carbonyls (C5-C9) were assigned as oxidation products of cycloalkanes. Therefore, we examined the correlations
 468 between m/z 69.070 and other cycloalkane fragments over the Arabian Gulf and Suez, where anthropogenic primary emissions
 469 were significant. As shown in Figure 5, m/z 83 was the most abundant fragment and it correlated better with m/z 69 than the other
 470 two masses, strongly supporting the presence of C6 cycloalkanes (methylcyclopentane and cyclohexane). The other two masses
 471 are distributed in two or three clusters, suggesting compositions of different cycloalkanes. M/z 43 and m/z 57 (fragments of C5
 472 cycloalkanes) had lower correlations with other fragments (not shown in the graph) as they are also fragments of other higher
 473 hydrocarbons. Thereby we could assign those unsaturated carbonyls as photochemical oxidation products (i.e. cyclic ketones or
 474 aldehydes) from their precursor cycloalkanes.



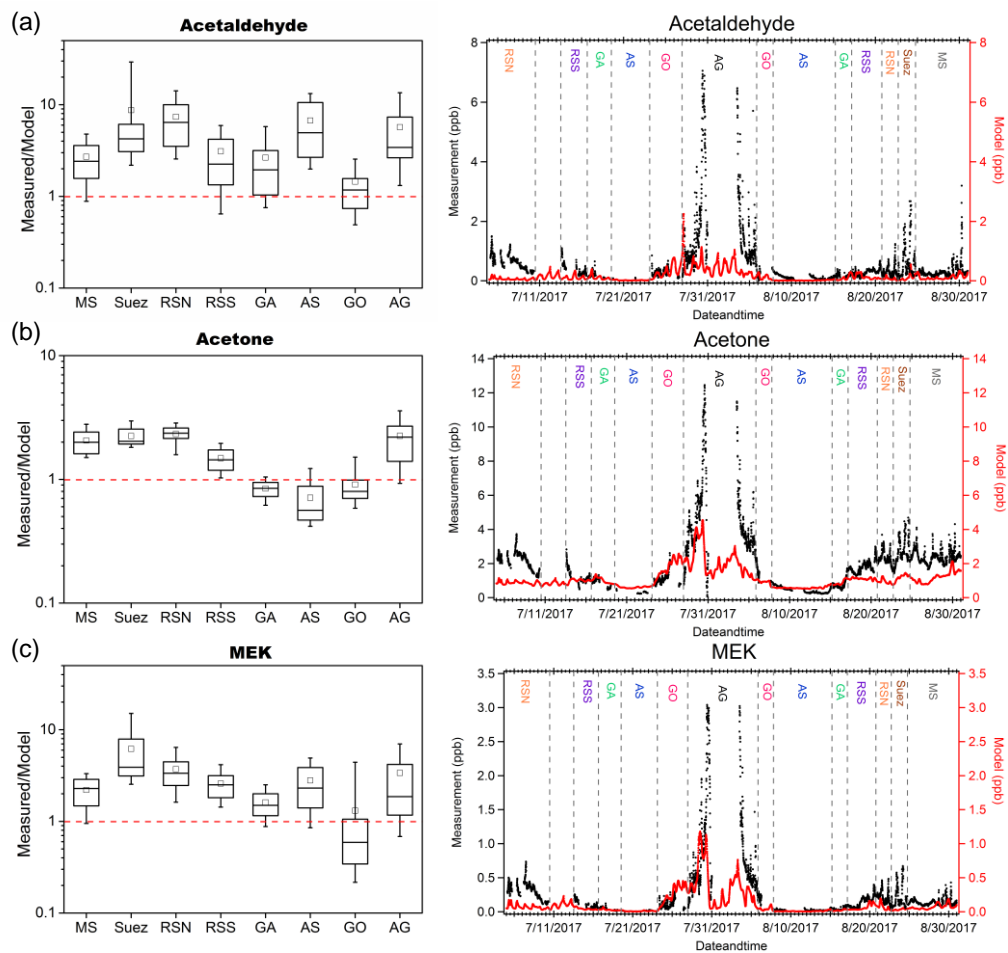
475
 476 Figure 5. Scatter plots of m/z 69.070 and other cycloalkane fragment masses over the (a) Arabian Gulf and (b) Suez region.
 477

478 As shown in Figure 2 and Table 1, C6 unsaturated carbonyls displayed higher mixing ratios than any other unsaturated carbonyls
 479 over the Arabian Gulf while C5 unsaturated carbonyl was slightly higher than C6 in Suez. Bourtsoukidis et al. (2019) derived
 480 enhancement ratio slopes from pentane isomers and established that the Arabian Gulf is dominated by oil and gas operations and
 481 that Suez is more influenced by ship emissions. Therefore, as the Arabian Gulf had much more active O&G activities than Suez,
 482 our findings agree with Koss et al. (2017) who showed that C6 unsaturated carbonyls should be more abundant than C5 carbonyls
 483 since more precursors for C6 unsaturated carbonyls are emitted from active oil fields. It is worth mentioning that in Figure 5 (b)
 484 one cluster at the bottom showed m/z 69.070 had no correlation with other three masses. Those points correspond to the time when
 485 the GC measured significant elevated isoprene while passing through the narrow Suez Canal where some vegetation (e.g. palms
 486 and some agriculture) was present close to shore, meaning m/z 69.070 during this period was isoprene. At the same time, m/z
 487 71.049 (C4 unsaturated carbonyl) increased from 20 ppt to 220 ppt. Isoprene oxidation products (MVK and methacrolein) were
 488 probably the major contribution to the C4 unsaturated carbonyls in this period. This also explains why C4 carbonyl dominated the
 489 distribution of unsaturated carbonyls over Suez.

490 In the other regions (especially more remote areas), the cyclic alkane fragmentation masses had much lower abundance, leading
491 to much less unsaturated carbonyls due to lack of precursors. Meanwhile, m/z 69.070 ($C_5H_8H^+$), m/z 83.086 ($C_6H_{10}H^+$) and m/z
492 97.101 ($C_7H_{12}H^+$) could also be fragmentations from corresponding aldehydes losing one water molecule as mentioned in section
493 2.3.3. Missing information of the chemical structure of unsaturated carbonyls and knowledge of their precursors, preclude detailed
494 investigation of the sources of large unsaturated carbonyls in these areas.

495 3.3 Model comparison of acetaldehyde, acetone and MEK

496 We compared our measurement results of acetaldehyde, acetone and MEK to those predicted by the global model “EMAC”
497 (ECHAM5/MESSy2 for Atmospheric Chemistry). From the results shown in Figure 6, the model predicted acetone much better
498 than acetaldehyde and MEK. In general, the model broadly captured the major features identified during the campaign such as
499 much higher levels of carbonyls mixing ratios over the Arabian Gulf and Suez and relatively low levels over the Arabian Sea. The
500 mean measurements-to-model ratios indicated that acetone was overestimated by a factor within 1.5 over the Arabian Sea, Gulf of
501 Aden and Gulf of Oman, and underestimated by a factor within 2.5 over the other regions. In contrast, the model underestimated
502 MEK within a factor of 4 over most of the regions except for the Gulf of Oman where MEK was overestimated (median values
503 were taken here as the mean values substantially deviated from the medians over Suez, Gulf of Oman and Arabian Gulf). The
504 model underestimation was most significant for acetaldehyde, which is underpredicted by a factor (median values) of more than 6
505 over the Red Sea North, ~ 4 over the Arabian Sea and Arabian Gulf and between 1 and 4 over other regions. A strong natural non-
506 methane hydrocarbon source from deep water in the Northern Red Sea was implemented in the model (Bourtsoukidis et al., 2020).
507 Although the model representation of acetaldehyde and other carbonyls was clearly improved after including the deep water source
508 of ethane and propane (Figure S4), the underestimation of acetaldehyde was still significant over the Red Sea North as shown in
509 Figure 6(a), indicating further missing sources. For acetaldehyde and MEK, the discrepancy was also significant over the Arabian
510 Sea where acetone was in contrast, overestimated. Since acetaldehyde had the biggest bias from the model prediction, we further
511 investigate the possible missing sources of acetaldehyde.



513

514 Figure 6. Measurement to model ratios (left) and time series (right) of measurements (in black) and model simulation (in red) of
 515 (a) acetaldehyde; (b) acetone; (c) MEK in each area. In each box plot, the box represents 25% to 75% of the data set with central
 516 line and square indicating the median value and the mean value respectively. The whiskers show data from 10% to 90%. The red
 517 dashed lines represent the 1:1 ratio.

518 3.4 Missing sources of acetaldehyde

519 In this section we investigate the following processes as potential sources of acetaldehyde: (1) production as an inlet artifact, (2)
 520 oceanic emission of acetaldehyde, (3) anthropogenic primary sources, (4) biomass burning sources, and (5) other possible
 521 secondary formation pathways.

522 3.4.1 Inlet artifact

523 Northway et al. (2004) and Apel et al. (2008) reported that heterogeneous reactions of unsaturated organic species with ozone on
524 the wall of the Teflon inlet can cause artifacts signal of acetaldehyde but not to acetone. During AQABA, the highest and the most
525 variable ozone mixing ratios were observed during the campaign over the Arabian Gulf (mean: 80 ± 34 ppb) and the Red Sea North
526 (66 ± 12 ppb), where a modest correlation was found between acetaldehyde and ozone over the Arabian Gulf ($r^2=0.54$) and no
527 significant correlation over the Red Sea North ($r^2=0.40$). However larger correlation coefficients were identified between ozone
528 and other carbonyls over the Arabian Gulf (see Figure S5), which suggests that the correlation was due to atmospheric
529 photochemical production rather than artifacts. Moreover, acetaldehyde was found to have a much worse correlation with ozone
530 during the nighttime compared to the correlation during the daytime over the Arabian Gulf (Figure 3b and c), which also indicates
531 that inlet generation of acetaldehyde was insignificant. Over other regions, especially the remote area (the Arabian Sea and Gulf
532 of Aden), ozone was relatively constant and low, with poor correlation with acetaldehyde mixing ratios. Although we cannot
533 completely exclude the possible existence of artifacts, the interference is likely to be insignificant in this dataset.

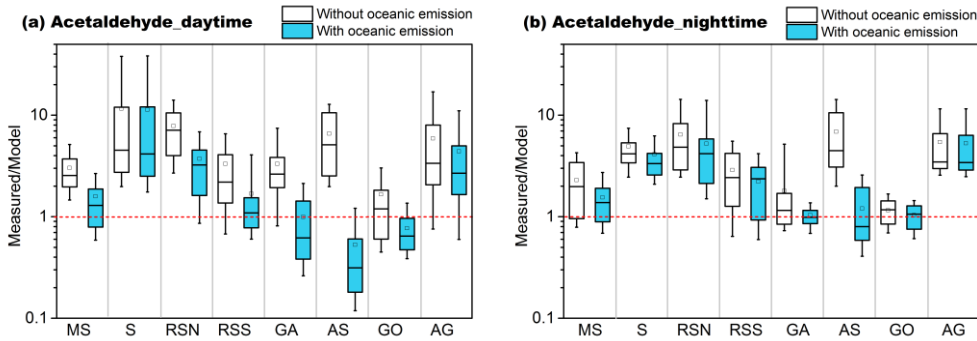
534 3.4.2 Oceanic emission

535 A bias between measured acetaldehyde and global model simulations has been observed in previous studies conducted in the
536 remote troposphere (Singh et al., 2003; Singh, 2004; Wang et al., 2019) and in the marine boundary layer (Read et al., 2012). The
537 aforementioned studies emphasized the potential importance of the sea water acting as a source of acetaldehyde emission via air-
538 sea exchange. No significant correlation was found between acetaldehyde and DMS, a marker of marine biogenic emission which
539 is produced by phytoplankton in seawater (Bates et al., 1992) (see Figure S6). This indicates that the direct biogenic acetaldehyde
540 emissions from the ocean are probably insufficient to explain the measured acetaldehyde. More likely, acetaldehyde and other
541 small carbonyl compounds can be formed in the sea especially in the surface microlayer (SML) via photodegradation of colored
542 dissolved organic matter (CDOM) (Kieber et al., 1990; Zhou and Mopper, 1997; Ciuraru et al., 2015). Zhou and Mopper (1997)
543 calculated the exchange direction of small carbonyls based on measurement results and identified that the net flux of acetaldehyde
544 was from sea to the air whereas formaldehyde was taken up by the sea. Sinha et al. (2007) characterized air-sea flux of several
545 VOCs in a mesocosm experiment and found that acetaldehyde emissions were in close correlation with light intensity ($r=0.7$). By
546 using a 3-D model, Millet et al. (2010) estimated the net oceanic emission of acetaldehyde to be as high as 57 Tg a^{-1} (in a global
547 total budget: 213 Tg a^{-1}), being the second largest global source. A similar approach was applied in a recent study done by Wang
548 et al. (2019), reporting the upper limit of the net ocean emission of acetaldehyde to be 34 Tg a^{-1} . Yang et al. (2014) quantified the
549 air-sea fluxes of several OVOCs over Atlantic Ocean by eddy covariance measurements, showing ocean is a net source for
550 acetaldehyde. Although Schlundt et al. (2017) reported uptake of acetaldehyde by the ocean from measurement-inferred fluxes in
551 western Pacific coastal regions, to our knowledge, there is no direct experimental evidence showing the ocean to be a sink for
552 acetaldehyde.

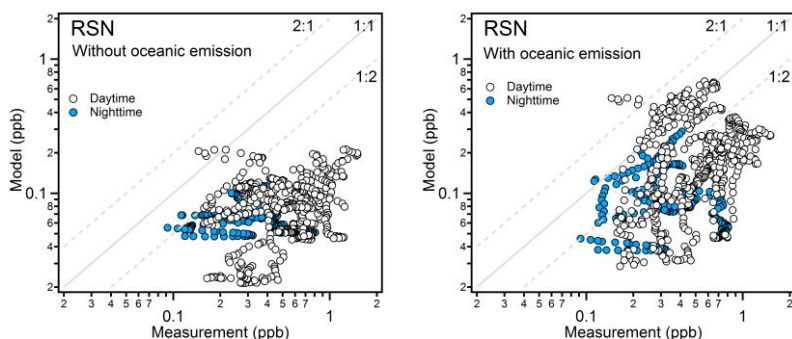
553 In order to test the importance of the oceanic emission of acetaldehyde, we implemented this source in EMAC model. The measured
554 sea water concentration of acetaldehyde was not available for the water area around the Arabian Peninsula. Wang et al. (2019)
555 estimated the global average acetaldehyde surface seawater concentrations of the ocean mixed layer using a satellite-based
556 approach similar to Millet et al. (2010), where the model estimation agreed well with limited reported measurements. From the
557 Wang et al. (2019) results, the averaged seawater concentration of acetaldehyde around Arabian Peninsula was generally much
558 higher from June to August. As the photodegradation of CDOM is highly dependent on sunlight, the air-sea submodel (Pozzer et
559 al., 2006) was augmented to include throughout the campaign a scaled acetaldehyde seawater concentration in the range of $0 \sim 50$
560 nM according to the solar radiation (Figure S7). With this approach, the average of acetaldehyde seawater concentration estimated
561 by the model is 13.4 nM , a reasonable level compared to predicted level by Wang et al. (2019).

562 After adding the oceanic source of acetaldehyde, the model estimation was significantly improved (Figure 7). As the oceanic source
 563 in the model is scaled according to the solar radiation, the measurement-to-model ratios were more strongly reduced during the
 564 day compared to the night. With oceanic emission included, the model underestimation was less significant, within a factor of 3
 565 during the day and 4 during the night over the Mediterranean Sea, Red Sea and Gulf of Aden. The most significant improvement
 566 was identified over the Red Sea North. As shown in Figure 8, the model had much better agreement with the measurement after
 567 adding the oceanic source. The scatter plots for other regions can be found in Figure S8. Over the Arabian Sea, the model
 568 significantly overestimated acetaldehyde mixing ratios, indicating the input sea water concentration of acetaldehyde might be too
 569 high. The SML layer starts to be effectively destroyed by the wave breaking when the wind speed exceeds than 8 m s^{-1} (Gantt
 570 et al., 2011). As the average wind speed over the Arabian Sea was the highest among the cruised areas ($8.1 \pm 2.4 \text{ m s}^{-1}$, Figure S1),
 571 less contribution from the CDOM photo degradation to acetaldehyde in the surface sea water would be expected. For the Suez
 572 region, due to the limited model resolution ($1.1^\circ \times 1.1^\circ$), little sea water was identified in the model, leading to negligible influence
 573 from the oceanic source.

574 Model underestimation of acetaldehyde especially over the Suez, Red Sea and Arabian Gulf is also likely to be related to the coarse
 575 model resolution ($\sim 1.1^\circ \times 1.1^\circ$) (Fischer et al., 2015). Where model grid points contain areas of land the higher and more variable
 576 terrestrial boundary layer height impacts the model prediction whereas the measurements may only be influenced by a shallower
 577 and more stable marine boundary layer.



578
 579 Figure 7. Acetaldehyde measurement to model ratios without the oceanic source (white boxes) and with the oceanic source (blue
 580 boxes) in the model during (a) daytime and (b) nighttime in different regions. The boxes represent 25% to 75% of the data set
 581 with the central line and square indicating the median and mean values, respectively. The whiskers show data from 10% to 90%.
 582 The red dashed lines represent the 1:1 ratio.



583

584 Figure 8. Observed and simulated mixing ratios of acetaldehyde over the Red Sea North without oceanic emission (left) and with
 585 oceanic emission (right). The data points are separated into day- and nighttime according to solar radiation.

586 **3.4.3 Anthropogenic primary sources**

587 Over the Arabian Gulf and Suez, the intensive photochemical production of carbonyls is apparent. Boursoukidis et al. (2020)
 588 compared measured hydrocarbons (ethane, propane, and butane) with the results from model simulations (the same model used in
 589 this study with the newly discovered deep water source implemented). The model was able to reproduce the measurement over
 590 most regions expect for some significant model underestimations in Suez and Arabian Gulf, in which local and small-scale
 591 emissions were difficult for the model to capture. Therefore, an underestimation of the precursor hydrocarbons, as well as those
 592 large alkanes, alkenes and cyclic hydrocarbons which were not measured (> C8) or included in the model (> C5) could be a reason
 593 for the model underestimation of acetaldehyde especially in polluted regions. In addition, as mentioned in the previous case studies,
 594 high ozone mixing ratios were observed over the Arabian Gulf especially during the nighttime. Ethene and propene were found to
 595 be significantly underestimated during the nighttime high ozone period by a factor over 10 (Figure S9), which indicates that the
 596 nighttime ozonolysis of alkenes could be another important source for acetaldehyde, formaldehyde and other carbonyls (Atkinson
 597 et al., 1995;Altshuller, 1993) in the Arabian Gulf.

598 Acetaldehyde, an oxygenated VOC, is not generally considered as an important primary emission from oil and gas field but instead
 599 a photochemical product of hydrocarbon oxidation (Yuan et al., 2014;Koss et al., 2015;Koss et al., 2017). In contrast, primary
 600 sources of formaldehyde from oil and gas production processes including both combustion and non-combustion process have been
 601 ascertained (Vaught, 1991). Le Baron and Stoeckenius (2015) concluded in their report of the Uinta Basin winter ozone study that
 602 besides formaldehyde, the other carbonyls were poorly understood in terms of their primary sources. Acetaldehyde and other
 603 carbonyls (aldehydes and ketones) have been reported as primary emissions from fossil fuel combustion including ship emissions
 604 (Reda et al., 2014;Xiao et al., 2018;Huang et al., 2018) and vehicle emissions (Nogueira et al., 2014;Erickson et al., 2014;Dong et
 605 al., 2014). A possible explanation for the measurement-model discrepancy is that the active petroleum industry located in the
 606 Arabian Gulf and intensive marine transportation in Suez are primary sources of acetaldehyde and other carbonyls which were not
 607 well constrained in the model. The Suez region, where the largest acetaldehyde discrepancy was identified, had a significant
 608 influence from biomass burning (see section 3.2.2). Biomass burning emissions are notoriously difficult to model as they are highly
 609 variable both in time and space. In this study, the model failed to reproduce the acetonitrile level with a range of only 40-50 ppt
 610 rather than 100-550 ppt measured over Suez. Thus, besides the possibility of seawater emission from the Gulf of Suez and the Suez
 611 Canal, the underestimated biomass burning source in the model over Suez, will lead to an underestimation of acetaldehyde as well
 612 as other carbonyl compounds in this region.

613 3.4.4 Other possible secondary formation pathways

614 Although the model estimation was generally improved with the addition of an oceanic source, the model to measured ratios still
615 varied over a wide range. As mentioned above, photodegradation of CDOM on the surface of seawater is a known source for
616 acetaldehyde although some studies focusing on real sea water samples did not observe clear diel cycles of seawater acetaldehyde
617 (Beale et al., 2013; Yang et al., 2014). Fast microbial oxidation could be a reason (Dixon et al., 2013) while other non-light driven
618 sources of acetaldehyde could be an alternative explanation. In a recent study, Zhou et al. (2014) reported enhanced gas-phase
619 carbonyl compounds including acetaldehyde during a laboratory experiment of ozone reacting with SML samples, indicating
620 acetaldehyde could also be produced under non-light driven heterogeneous oxidation. Wang et al. (2019) ventured a hypothetical
621 source that organic aerosol can be an extra source for unattributed acetaldehyde in the free troposphere through light-driven
622 production and ozonolysis. However, since the yield of acetaldehyde from such reactions is unknown, large uncertainties remain.
623 Previous studies have shown that the organic matter fraction was highest in smaller sea spray aerosols and that the aerosols contain
624 both saturated and unsaturated fatty acids originating from the seawater surface (i.e. SML) (Mochida et al., 2002; Cochran et al.,
625 2016). Thus, for the AQABA campaign, both photodegradation and heterogeneous oxidation could occur on the surface of sea
626 spray and pollution associated aerosols, even over remote open ocean therefore being an extra source of acetaldehyde and other
627 carbonyl compounds.

628 Another acetaldehyde formation pathway reported is gas-phase photolysis of pyruvic acid (Eger et al., 2019b; Reed Harris et al.,
629 2016), a compound mainly of biogenic origin. Pyruvic acid has been also observed in seawater (Kieber and Mopper, 1987; Zhou
630 and Mopper, 1997) and was found up to 50 nM in the surface water of eastern Pacific Ocean (Steinberg and Bada, 1984), while
631 acetaldehyde was not the major product of aqueous-phase photolysis of pyruvic acid (Griffith et al., 2013). Zhou and Mopper
632 (1997) pointed out that the net exchange direction for pyruvic acid is expected to be from the air to the sea due to high solubility,
633 with Henry's law constant of $3.1 \times 10^3 \text{ mol m}^{-3} \text{ Pa}^{-1}$ (Sander, 2015). Moreover, partitioning to aerosols could be an important sink
634 for pyruvic acid (Reed et al., 2014; Griffith et al., 2013): an increasing concentration trend of pyruvic acid was observed in marine
635 aerosols over western North Pacific Ocean (Boreddy et al., 2017). Therefore, due to limited terrestrial biogenic sources of pyruvic
636 acid for AQABA campaign, gas-phase level of pyruvic acid was expected to be low. Limited studies reported pyruvic acid level
637 in marine boundary layer, Baboukas et al. (2000) measured 1.1 ± 1.0 ppt of pyruvic acid above Atlantic Ocean. Pyruvic acid was
638 measured by Jardine et al. (2010) using a PTR-MS at m/z 89 in a forested environment. For the AQABA PTR-ToF-MS data set,
639 enhanced signals were observed at m/z 89.024 with the mean mixing ratio of 35-110 ppt over different regions (Table S4), which
640 is much more abundant than reported pyruvic acid levels by Baboukas et al. (2000). This might be due to the uncertainty associated
641 with the theoretical methods of quantification used here or the presence of isomeric compounds on that mass, since pyruvic acid
642 was not calibrated with the standard. Even if we assume the m/z 89.024 to be entirely pyruvic acid, with 60% yield of acetaldehyde
643 via photolysis (IUPAC, 2019), it gave maximum 13 ppt of acetaldehyde over Arabian Gulf, 5-9 ppt over other regions, which were
644 only 0.8% - 6% of the mean mixing ratios (Table S4). Detailed information of the calculation can be found in the Supporting
645 Information. Therefore, we conclude that the contribution from the photolysis of pyruvic acid is not an important source for the
646 unattributed acetaldehyde during the AQABA campaign.

647 4 Summary and Conclusion

648 Observations of carbonyl compounds around the Arabian Peninsula were investigated in terms of mixing ratios abundance over
649 different areas. Aliphatic carbonyl compounds were generally more abundant than the unsaturated and aromatic carbonyl
650 compounds, and were dominated by low-molecular-weight compounds (carbon number less than five). Aliphatic carbonyl

651 compounds were found at the highest mixing ratios over the Arabian Gulf followed by the Suez region, while the lowest mixing
652 ratios were observed over the Arabian Sea and the Gulf of Aden. Over the Mediterranean Sea, aliphatic carbonyls were low except
653 for acetone that was much higher compared to the levels observed over clean remote areas (i.e. Arabian Sea). The atmospheric
654 composition over the Red Sea showed obvious differences between the northern and the southern part, with higher mixing ratios
655 in the north. Similar region-dependent distributions were observed for unsaturated and aromatic carbonyls. Generally, the mixing
656 ratios of aromatic carbonyl compounds decreased as the carbon number increased. Particularly over the Suez region, benzaldehyde
657 (C7 aromatic carbonyls) was much more abundant than other aromatic carbonyls, indicating direct sources as well as abundant
658 oxidation precursors. For unsaturated carbonyl compounds, C5 and C6 carbonyl compounds dominated the mixing ratio
659 distribution, while the air chemistry highly depends on the chemical structure assignment of those masses.

660 Further case studies showed that the carbonyl compounds were highly correlated to the high ozone levels during daytime over the
661 Arabian Gulf while the air chemistry in Suez region was strongly influenced by regional biomass burning. Due to the unexpectedly
662 high loading of m/z 69 (usually assigned as isoprene) observed in highly polluted regions, we further identified the correlations
663 between m/z 69 and other fragmentation masses of cycloalkanes according to previous studies conducted in oil and gas regions
664 (Warneke et al., 2014; Yuan et al., 2014; Koss et al., 2017). The high correlations among fragments implied the existence of
665 cycloalkanes in the polluted regions, which could be further oxidized to unsaturated carbonyl compounds (cyclic ketones or
666 aldehydes).

667 As acetaldehyde was identified as having important additional sources, we further compared the measurements of major carbonyl
668 species (acetaldehyde, acetone and MEK) with a comprehensive global atmospheric chemistry model (EMAC). Acetaldehyde was
669 found to have the highest discrepancy between the observations and model simulations, with the simulated values to be lower up
670 to a factor of 10. By adding an oceanic source of acetaldehyde produced via light-driven photodegradation of CDOM in the
671 seawater, the model estimation improved significantly, especially over the Red Sea North. With the oceanic source added, modelled
672 acetaldehyde became slightly overestimated in clean regions, suggesting that the emission rate employed represents an upper limit.
673 The results indicate that the ocean plays an important role in the atmospheric acetaldehyde budget, under both clean and polluted
674 conditions. The underestimated acetaldehyde in the model is significant as it will influence the atmospheric budget of e.g. PAN.
675 As shown in Figure 1, multiple sources and formation pathways need to be considered to better understand the atmospheric budget
676 of acetaldehyde. Additional laboratory experiments and field measurements are necessary in order to verify all possible
677 atmospheric formation mechanisms and to improve model simulations.

678 **Data availability.**

679 [The data used in this study are available to all scientists agreeing to the AQABA protocol at](https://doi.org/10.5281/zenodo.3974228)~~Data will be made available via:~~
680 <http://doi.org/10.5281/zenodo.3974228> (Wang et al., 2020). <https://edmond.mpdl.mpg.de/imeji/>

681 **Author contributions.**

682 AE and CS performed PTR-ToF-MS measurement and preliminary data processing. NW conducted data analysis and drafted the
683 article. AP performed EMAC model simulation. EB and LE are responsible for NMHC measurements and data. DD, BH and HF
684 provided formaldehyde data. Ozone and actinic flux data were contributed by JS and JNC. Methane and carbon monoxide data
685 were provided by JP. JL designed and realized the campaign. JW supervised the study. All authors contributed to editing the draft
686 and approved the submitted version.

687 **Competing interest.**

688 The authors declare that they have no conflict of interest.

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