

Trace Organic Gas Analyzer Time-of-Flight mass spectrometer (TOGA-TOF) system for airborne observations of formaldehyde

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41 **Abstract**

42 Formaldehyde (HCHO) is a ubiquitous atmospheric constituent, originating from primary
43 emissions (natural and anthropogenic) and secondary production via the oxidation of volatile
44 organic compounds (VOCs). In addition to being a regulated pollutant, HCHO is a key species
45 used as a tracer of recent photochemical activity due to its short atmospheric lifetime and its role
46 as a source of HO_x radicals. Given its diverse sources and high spatial variability, HCHO is
47 challenging to represent accurately in chemical transport models, often resulting in significant
48 discrepancies with observations. Airborne *in situ* measurements of HCHO, especially when
49 combined with VOC precursor data, offer valuable insights into its atmospheric distributions for
50 evaluating models. Here, we present HCHO observations from the NSF NCAR Trace Organic
51 Gas Analyzer with Time-of-Flight mass spectrometer (TOGA-TOF), deployed during the 2019
52 Fire Influence on Regional to Global Environments and Air Quality (FIREX-AQ) campaign.
53 While most HCHO instruments target at most a few selected species for measurement, the TOGA-
54 TOF employs a rapid gas chromatography-mass spectrometry (GC/MS) technique and provides
55 discrete VOC measurements—including >100 C₁–C₁₀ species—at a time resolution of less than
56 2 minutes. We compare TOGA-TOF HCHO data to measurements from three 1 Hz instruments
57 aboard the NASA DC-8: the Compact Atmospheric Multi-species Spectrometer (CAMS), the In
58 Situ Airborne Formaldehyde (ISAF) instrument, and a proton-transfer-reaction time-of-flight
59 mass spectrometer (PTR-ToF-MS). The wide dynamic range of observed HCHO concentrations
60 (from < 100 ppt to ~100 ppb) during FIREX-AQ enabled a robust intercomparison. TOGA-TOF
61 HCHO agreed well with CAMS (slope = 1.1), with similar agreement with the PTR-ToF-MS,
62 while larger discrepancies were observed with ISAF (slope = 1.5), likely due to differences in
63 calibrations. Normalized excess mixing ratios (NEMRs) of HCHO relative to CO in wildfire
64 plumes exhibited consistent trends with plume age across instruments. These findings highlight
65 the TOGA-TOF's capability for highly sensitive and accurate airborne HCHO measurements.

66 **1. Introduction**

67 Formaldehyde (HCHO) is one of the most ubiquitous non-methane volatile organic
68 compounds (VOCs) in the atmosphere. In addition to being an EPA regulated pollutant (National
69 Research Council (US) Committee on Toxicology, 1980), HCHO is a key reactive compound in

70 understanding recent photochemistry in the atmosphere. HCHO is an oxidation product of a wide
71 range of VOCs, has a relatively short lifetime, and is an important source of HO_x (OH + HO₂),
72 therefore affecting NO_x-VOC-O₃ chemistry (Seinfeld and Pandis, 2006). Globally, the dominant
73 source of HCHO is secondary production from VOC oxidation (Luecken et al., 2012; Stavroukou
74 et al., 2009). Tropospheric background levels of HCHO are maintained through methane (CH₄)
75 (Stavroukou et al., 2009) and isoprene (C₅H₈) (Curci et al., 2010; Luecken et al., 2018; Palmer et
76 al., 2003; Wolfe et al., 2016) oxidation. Direct emissions of HCHO can also be important locally;
77 sources include vehicle emissions (Possanzini et al., 2002; Sagebiel et al., 1996; Viskari et al.,
78 2000), biomass burning (Lee et al., 1997; Zhang et al., 2013), and fossil fuel industries (Fried et
79 al., 2020; Green et al., 2021; Pfister et al., 2019). In the atmosphere, HCHO is mainly lost through
80 photolysis, reaction with OH, and deposition, resulting in daytime atmospheric lifetimes of 1 to 9
81 hours (1–3 hours midday).

82 As an important proxy for understanding recent photochemistry, HCHO observations have
83 been used to indicate ozone sensitivity regimes (VOC- or NO_x-limited) (Duncan et al., 2010;
84 Martin et al., 2004; Nussbaumer et al., 2021; Schroeder et al., 2017; Souri et al., 2020, 2023) as
85 well as top-down estimations of isoprene (Cao et al., 2018; Kaiser et al., 2018; Palmer et al., 2003;
86 Wolfe et al., 2016; Zhu et al., 2016) and anthropogenic VOC (Cao et al., 2018; Kwon et al., 2021)
87 emissions. While satellite retrievals of HCHO column densities have been widely used in these
88 methods, challenges remain in capturing the spatial (vertical and horizontal) representation of
89 HCHO over the wide mixing ratio (MR) range observed in the troposphere; HCHO MRs can vary
90 less from than 20 parts per trillion (ppt) in remote regions, 1–20 parts per billion (ppb) in urban
91 environments, and > 100 ppb in fire plumes as observed from aircraft. These challenges in HCHO
92 satellite observations have been driven by uncertainties in retrieval methods (Anderson et al., 2017;

93 Liao et al., 2025; Souri et al., 2023; Stavrakou et al., 2009; Zhu et al., 2016), instrument sensitivity
94 (Souri et al., 2023), and coarse pixel sizes (Kwon et al., 2021; Souri et al., 2023). Uncertainties in
95 satellite observations increase in environments with low HCHO, such as regions with weak
96 biogenic or anthropogenic sources (Souri et al., 2023). Three-dimensional chemical transport
97 models may also fail to reproduce observed HCHO levels due to inaccurate emission inventories
98 of direct HCHO emissions or its VOC precursors (Green et al., 2021; Jaeglé et al., 2018; Luecken
99 et al., 2012, 2018; Warneke et al., 2007), uncertainties in chemical mechanisms (Luecken et al.,
100 2012), and coarse grid size. High quality *in situ* HCHO observations during airborne field
101 campaigns can provide useful information to validate satellite observations and evaluate chemical
102 transport models (Anderson et al., 2017; Chan Miller et al., 2017; Schroeder et al., 2017; Souri et
103 al., 2023; Zhu et al., 2016). For example, recent studies on HCHO intercomparisons between
104 satellite and *in situ* airborne observations showed that satellite HCHO retrievals can have a
105 systematic low bias of -20 to -51% (Anderson et al., 2017; Zhu et al., 2016), which can lead to
106 underestimation of VOC emissions. These systematic biases in satellite observations can be
107 corrected using high-quality airborne *in situ* observations of HCHO (Chan Miller et al., 2017; Zhu
108 et al., 2016, 2017).

109 Biomass burning produces large amounts of HCHO both from primary emissions and
110 secondary production. During a biomass burning event, a highly complex mixture of VOCs (Akagi
111 et al., 2011; Gilman et al., 2015; Gkatzelis et al., 2024; Koss et al., 2018; Travis et al., 2023;
112 Yokelson et al., 2013a) is produced from the fire source, which subsequently undergoes complex
113 chemical (i.e., reaction with oxidants and photolysis) and physical (i.e., dilution and cooling)
114 transformations downwind of the source (Decker et al., 2021; Hornbrook et al., 2011; Robinson et
115 al., 2021; Wang et al., 2021). For the recent joint NOAA/NASA Fire Influence on Regional to

116 Global Environments and Air Quality (FIREX-AQ) mission, Wang et al. (2021) used a high-
117 resolution large eddy simulation (LES) to show the chemical complexity of a fresh wildfire plume.
118 The levels and types of predominant oxidants, which are dependent on the historic exposure to
119 light, varies with the evolution of a plume, affecting the chemistry and fate of trace gases and
120 aerosols (Decker et al., 2021; Robinson et al., 2021; Wang et al., 2021). Liao et al. (2021) showed
121 that primary emissions of HCHO dominate near the fire source but secondary production from
122 VOC oxidation becomes important (up to ~ 60% after a few hours of physical age) as the fire ages.
123 As reported by Travis et al. (2023), the MRs of different types of VOCs emitted in crop and
124 prescribed fire plumes during FIREX-AQ depends on fuel types and fire states. The physical and
125 chemical complexities in fire plumes ensure that HCHO MRs are highly dynamic presenting
126 challenges for satellite retrievals (Stavrakou et al., 2009) and models to accurately represent local
127 and regional distributions of HCHO.

128 In this study, we present the NSF NCAR Trace Organic Gas Analyzer Time-of-Flight mass
129 spectrometer (TOGA-TOF) as a fast online GC/MS (gas chromatography-mass spectrometry)
130 technique for airborne observations of HCHO. Studies on improved techniques to detect HCHO
131 have been ongoing, attesting to its analytical challenges (Gilpin et al., 1997; Hopkins et al., 2003;
132 Hunter et al., 1999; Rice and Quay, 2006). While most currently available instrumental techniques
133 are specialized in detecting HCHO, the TOGA-TOF routinely measures over a hundred VOCs
134 using a non-reactive concentration technique, with the capability to analyze untargeted
135 compounds. Thus, TOGA-TOF is a unique airborne standalone instrument for the identification
136 and quantification of a wide range of C_1 – C_{10} VOCs, including formaldehyde, in complex ambient
137 air samples. The NOAA/NASA FIREX-AQ campaign (July to September 2019) was the first
138 campaign in which the TOGA-TOF was deployed for airborne measurements. During FIREX-AQ,

139 we had the opportunity to measure alongside two state-of-the-art HCHO measurement systems
140 aboard the NASA DC-8 during the entire FIREX-AQ campaign: the NASA In Situ Airborne
141 Formaldehyde (ISAF) instrument (Cazorla et al., 2015) and the University of Colorado Compact
142 Atmospheric Multi-species Spectrometer (CAMS) (Richter et al., 2015). The University of Oslo
143 Proton-Transfer Reaction Time-of-Flight Mass Spectrometer (UiO PTR-ToF-MS) (Müller et al.,
144 2014) measured HCHO for select time periods as well. For this study, we present the TOGA-TOF
145 HCHO data and focus on intercomparisons with CAMS and ISAF but also include comparisons
146 with the UiO PTR-ToF-MS. The FIREX-AQ dataset is an excellent testbed to compare a wide
147 range of HCHO mixing ratios (below 100 ppt to ~100 ppb) in a spatially variable ambient
148 atmosphere. The technique used in the TOGA-TOF could, in principle, be implemented in many
149 existing online GC/MS systems for semi-routine measurements of HCHO.

150 **2. Methods**

151 The sample preconcentration system in the TOGA-TOF is similar to the earlier generation
152 TOGA instrument with a quadrupole mass analyzer, which has been described in the literature
153 (Apel et al., 2015). The TOGA-TOF has been extensively characterized in the lab prior to and post
154 FIREX-AQ. Here, we describe the basic operation techniques of the TOGA-TOF system (**Sect.**
155 **2.1**), summarize the HCHO measurements during FIREX-AQ (**Sect. 2.2**), calculate the weighted
156 averages for direct comparisons between instruments with different sampling time resolution
157 (**Sect. 2.3**), and derive normalized excess mixing ratios (NEMR) of HCHO from selected plume
158 transects (**Sect. 2.4**).

159 **2.1 Description of the NSF NCAR Trace Organic Gas Analyzer with Time-of-Flight Mass**
160 **Spectrometer (TOGA-TOF)**

161 The TOGA-TOF hardware consists of a cryogenic preconcentration system, gas
162 chromatograph (GC), time-of-flight (TOF) mass analyzer (TOFWERK, Switzerland), electronics
163 box, clean air generator and on-board calibration system (CAG), and pump box. All major
164 components except the TOF mass analyzer are custom built. Typical preconcentration and GC
165 analysis sequences for the system are shown in **Fig. 1** and the flows for each sequence are shown
166 in **Fig. 2**. The GC analysis, including temperature ramp and cool and sample preconcentration
167 steps, occur simultaneously, with the full cycle duration of both processes time synced (**Fig. 1**) to
168 less than 2 min. During FIREX-AQ, the runtime for one full cycle was 105 seconds including 33
169 seconds of sample collection (i.e., “sampling”). The preconcentration system consists of three
170 cryogenically-cooled traps as described by Apel et al. (2015): (1) a water trap for removal of water
171 in ambient sampled air, (2) a sample trap for enrichment of targeted VOCs in ambient air, and (3)
172 a cryofocusing trap to refocus the analytes prior to injection into the GC. Corresponding to each
173 temperature setpoint described in **Fig. 1**, the cooling of each trap is achieved by metering the flow
174 of gas-phase N₂ generated from the headspace of the TOGA-TOF Dewar through coils at the
175 bottom of the dewar that were immersed in liquid N₂ (Apel et al., 2015). To heat the traps, the cold
176 N₂ gas flow was first shut off and followed by PID controlled flow of electricity, via a
177 WATLOW™ (Anafaze MLS300) temperature controller, through resistive wires wrapped around
178 the traps. The sample trap is packed with glass wool to allow greater surface area for the
179 enrichment of VOCs when cooling the trap to -140°C (**Fig. 1**), while allowing bulk gases such as
180 oxygen and nitrogen to pass through unretained. Both the water trap (inner diameter (I.D.) = 0.04
181 in.) and cryofocusing trap (I.D. = 0.021 in.) are open tubes.

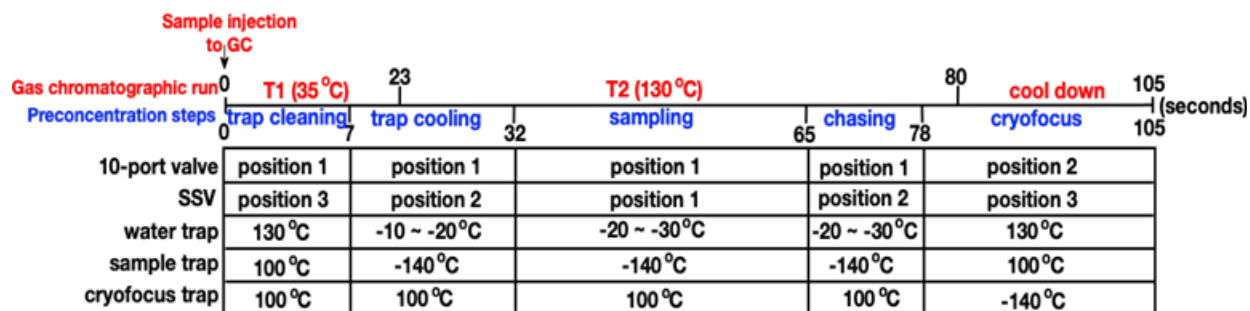


Figure 1. TOGA-TOF operational sequences for the gas chromatograph temperature program (red) and the pre-concentration steps (blue). The position of the valves (10-port valve and stream selector valve; SSV) and temperatures of each trap are shown for each step. The duration of each sequence corresponds to the normal sampling mode timing during FIREX-AQ.

182

183 For chromatography, a Restek MXT-624 column (I.D. = 0.18 mm, length = 8 m) is used
 184 with helium as the carrier gas. The GC column has two temperature set points with temperatures
 185 ranging between 35 °C and 130 °C (**Fig. 1**), also controlled by the Watlow, allowing gas-phase C₁
 186 to C₁₀ VOCs to sequentially elute from the column. This allows the separation of VOCs, including
 187 many structural isomers with the same chemical formula, based on their differences in interaction
 188 with the stationary phase of the column. For detection of VOCs, the GC is coupled to a high-
 189 resolution electron ionization time-of-flight mass spectrometer (HR EI-TOF-MS; TOFWERK),
 190 using 70 eV ionization resulting in classical electron ionization (EI) spectra of the compounds.
 191 This methodology allows routine quantification of well over 100 VOCs with the ability to further
 192 analyze additional C₁ to C₁₀ compounds within the detectable range. Fluctuations in sensitivity are
 193 corrected by normalizing with ambient levels of tetrachloromethane (CCl₄) based on the
 194 recommendations in Karbiwnyk et al. (2003). Background corrections are made by system
 195 generated ultra-pure helium (“He mode”) and zero-air blanks and then subtracting average blank
 196 peak areas from ambient samples. For all surfaces that the sample contacts, Restek Sulfinert[®] steel
 197 tubing is used to minimize interactions. Data processing is performed using Tofware (Aerodyne
 198 Research, Inc., Billerica, MA) for high resolution peak fitting and TERN-in-IGOR (Claflin et al.,

199 2021; Isaacman-VanWertz et al., 2017; Lerner et al., 2017) (Aerodyne Research, Inc.) for
 200 chromatographic peak analysis.

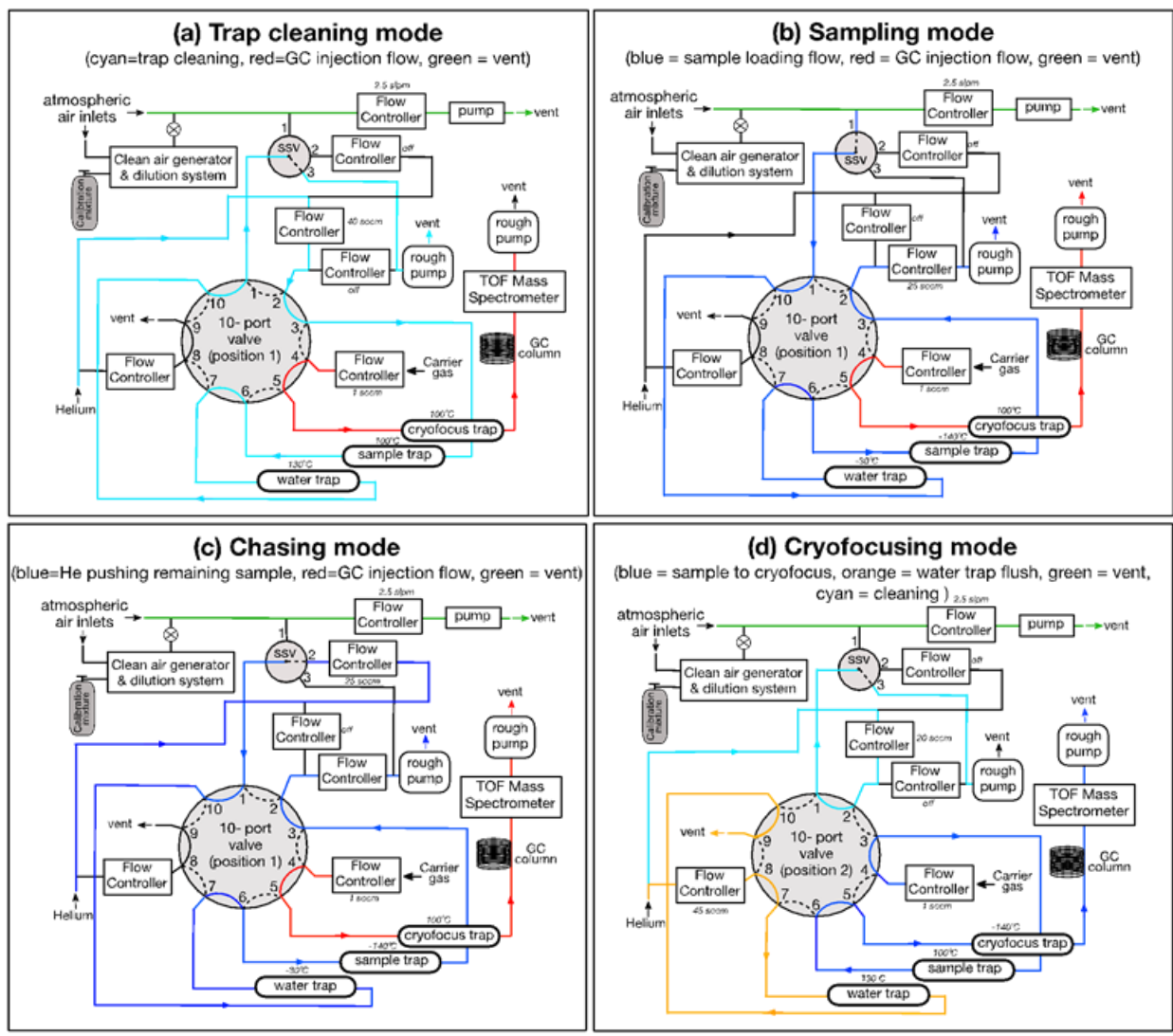


Figure 2. Simplified flow diagram of TOGA-TOF showing flows and temperatures of each trap during (a) trap cleaning mode, (b) sampling mode, (c) chasing mode, and (d) cryofocusing mode, corresponding to the sequence in Fig. 1.

201

202 **2.2 Description of HCHO and CO measurements during FIREX-AQ**

203

In this section, we describe the TOGA-TOF HCHO measurements during FIREX-AQ

204

(July–September 2019). FIREX-AQ (Warneke et al., 2023;

205

<https://asdc.larc.nasa.gov/project/FIREX-AQ>) focused on the emissions of trace gases and

206 particles from wildland, prescribed, and agricultural fires in the United States. The NASA DC-8
207 flights during FIREX-AQ were based out of Boise, ID, mainly focusing on Western U.S. fires, and
208 Salina, KS, focusing on southeastern U.S. agricultural fires (flight tracks shown in **Fig. S1**). Details
209 on the field campaign may be found in Warneke et al. (2023).

210 As described in **Sect. 2.1**, the TOGA-TOF uses electron ionization for the detection of
211 VOCs. The EI of HCHO results in a spectrum of its parent ion (m/z 30) and major fragments (m/z
212 28 and 29) (**Fig. S2a**). **Figure S2b** shows the chromatographic signals of CH_2O^+ (m/z 30; exact
213 mass 30.010016 amu) and CHO^+ (m/z 29; exact mass 29.002191 amu) eluting at ~ 20 s. The high
214 resolution (HR)-TOF allows detection of highly resolved ions and the Tofware peak fitting
215 analysis shows separate parent and dominant fragments from other adjacent peaks (**Fig. S2c**).
216 HCHO calibration was carried out using a HCHO in N_2 gas-phase mixture contained in a treated
217 aluminum cylinder (Apel-Riemer Environmental, Inc.), diluted with ultra-high purity N_2 . The
218 stated HCHO concentration of the calibration mixture cylinder from the company was 1.69 ppmv.
219 The concentration was checked using three independent methods within a two-month period using:
220 (1) Fourier transform infrared spectroscopy (FTIR – NSF NCAR ACOM direct absorption using
221 the Beer-Lambert law, 1.55 ppmv), (2) Mid-IR – University of Colorado – also direct absorption
222 employing the CAMS instrument (1.65 ppm), and (3) an EPA 2,4-dinitrophenylhydrazine (DNPH)
223 cartridge method (1.64 ppmv). For methods (1) and (2), quantification of HCHO relies on the
224 absolute IR cross section and thus does not require an external standard. Therefore, these can be
225 regarded as fully independent reference methods. For the calibration of TOGA-TOF we used the
226 averaged value of 1.61 ppmv from the three quantification methods, and the resulting calibration
227 curve is shown in **Fig. S3**. The TOGA-TOF HCHO lower limit of detection (LLOD) during
228 FIREX-AQ was 20 ppt, and the HCHO measurement had an uncertainty of 35% based on the

229 variability of repeat calibrations using a 1.6 ppm (parts per million) HCHO in N₂ standard mixture
230 and a 90 ppb HCHO in N₂ standard mixture (both Apel-Reimer, Inc.), dynamically diluted to MRs
231 ranging between 0.15 and 100 ppb using (1) the CAG (i.e., catalytically-scrubbed ambient air), (2)
232 dry N₂, and (3) humidified N₂. No humidity dependence was observed.

233 HCHO mixing ratios from the TOGA-TOF were compared to 1 Hz HCHO measurements
234 from the ISAF, CAMS, and UiO PTR-ToF-MS instruments onboard the DC-8. The ISAF
235 instrument was operated by NASA and the details on the instrumental technique are described in
236 Cazorla et al. (2015). Briefly, the ISAF instrument uses a laser-induced fluorescence technique to
237 quantify HCHO using the fluorescence resulting from excitation of HCHO at 353 nm with a
238 tunable UV laser. Backgrounds were determined from the offline position 0.005 nm away from
239 the peak. Calibrations were made using a compressed HCHO gas cylinder (Air Liquide, 584 ppbv)
240 quantified by FTIR (Liao et al., 2021). ISAF was operated at a native sampling rate of 10 Hz and
241 archived at 10 Hz and 1 Hz. We used the 1 Hz data for analysis. The reported LLOD of HCHO
242 measured by ISAF is 30 ppt for the 1 Hz data at a signal/noise = 1 and the accuracy (systematic
243 uncertainty) of the ISAF measurements is estimated to be 10% + 10 ppt. More details on the ISAF
244 HCHO measurement during FIREX-AQ can be found in Liao et al. (2021).

245 The CAMS was operated by the University of Colorado and a detailed description of the
246 system can be found in the work of Richter et al. (2015). The CAMS uses a tunable IR laser source
247 for generation of mid-IR light for the detection of HCHO based on absorption. Background spectra
248 were collected during flights using zero air from a scrubber and were subtracted from the ambient
249 spectra. Calibrations were carried out by using a 4.9 ppm HCHO standard in N₂, quantified by
250 direct absorption using the Beer-Lambert law. The reported uncertainty of the FIREX-AQ HCHO
251 CAMS measurements is 6% and the campaign-averaged LLOD is 110 ppt for 1 Hz measurements

252 at a signal/noise = 1 during FIREX-AQ. However, during previous airborne missions (e.g.,
253 KORUS-AQ; Fried et al., 2020), LLODs ranging from 28 to 80 ppt have been reported. The
254 degraded performance during FIREX-AQ resulted from dirty multipass cell mirrors.

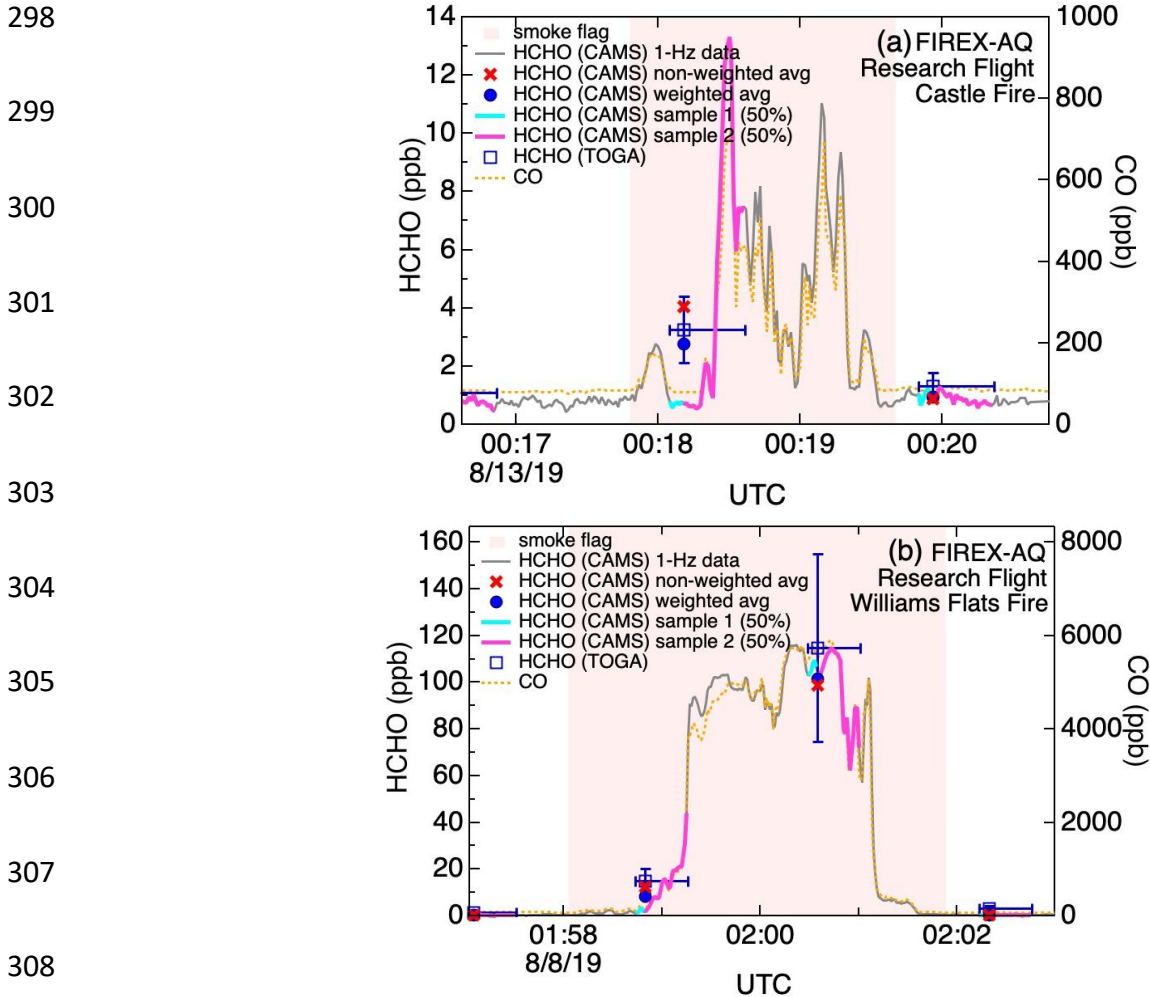
255 The UiO PTR-ToF-MS (Müller et al., 2014) was operated by the University of Oslo with
256 a focus on measuring NH₃ during FIREX-AQ, although HCHO and a number of additional VOCs
257 were reported for select flights (i.e., 25 July and 3 August). The PTR-ToF-MS uses hydronium
258 ions (H₃O⁺) generated from water vapor in a glow discharge-drift-tube to detect VOCs through
259 nondissociative proton transfer reactions. HCHO calibrations were carried out using a gas-phase
260 HCHO in N₂ cylinder (Apel-Riemer Environmental, Inc., 9.6 ppm HCHO; ± 5%). This calibration
261 was checked post-mission by generating test atmospheres with five different HCHO mixing ratios
262 (145–1035 ppb) in a 250 L laboratory chamber. The chamber was equipped with a White cell type
263 multiple reflection mirror system with a 120 m optical path length for online detection with a
264 Bruker IFS 66v/S FTIR instrument. The two methods were in excellent agreement, with the PTR-
265 ToF-MS reporting a volume mixing ratios less than 3% higher than the FTIR. During the two flight
266 days (25 July and 3 August) the UiO PTR-ToF-MS measured HCHO; the slope of the retrieved
267 ambient HCHO relative to that from CAMS was 1.03.

268 Normalized excess mixing ratios (NEMRs) of HCHO to CO are calculated as described in
269 **Sect. 2.4**. CO measurements were made onboard the DC-8 using the NOAA Los Gatos Research
270 (LGR) CO instrument and the NASA Differential Absorption Carbon monOxide Measurement
271 (DACOM) instruments. The LGR instrument (Bourgeois et al., 2022) uses an infrared laser off-
272 axis integrated-cavity-output spectroscopy technique with a time response of 1 Hz. The DACOM
273 instrument uses a differential absorption technique with an infrared tunable diode laser detecting
274 CO at the 4.7 μm wavelength, giving a data frequency of 5 Hz and measurement uncertainty of

275 2% (Sachse et al., 1987). The two CO instruments were in excellent agreement with the coefficient
276 of determination $r^2 = 0.97$, and a slope of 0.97 during the entirety of the campaign (**Fig. S4**), with
277 the reported LGR CO consistently 2.2% below that of DACOM.

278 **2.3 Averaging methods for comparison of HCHO between different instruments**

279 Data averaging across various sampling time periods is often carried out for direct
280 comparison between instrumental techniques with different sample integration times. **Figure 3**
281 shows HCHO measured by TOGA-TOF and CAMS with LGR CO measurements during plume
282 transects from the Castle Fire (**Fig. 3a**) and the Williams Flats Fire (**Fig. 3b**). As described in **Sect.**
283 **2.1**, during FIREX-AQ, the TOGA-TOF took discrete samples for 33 seconds in a 105 second
284 cycle period; therefore, subsections of each plume transect were sampled for analysis (**Fig. 3**). To
285 directly compare the discrete TOGA-TOF HCHO sampling data, CAMS, ISAF, and UiO PTR-
286 ToF-MS 1 Hz HCHO data are averaged over the time during which the TOGA-TOF sampling
287 occurred. Here we use two averaging methods for HCHO comparison: (1) normal averaging and
288 (2) volume-weighted averaging. The normal averaging is performed by taking the arithmetic mean
289 of the 1 Hz source data measured between the start and stop times of TOGA-TOF sampling. Below
290 LLOD values of the source data are included in the averaging. A minimum of at least one valid
291 (i.e., measured value or below LLOD) 1-Hz source data point during the TOGA-TOF sampling
292 period is allowed for the analysis to produce a non-missing merged value. This algorithm is how
293 the NASA online merging tool generates the TOGA.DC8_MRG and is publicly available on the
294 NASA Langley Research Center (LaRC) Airborne Science Data repository at [https://www-
295 air.larc.nasa.gov/cgi-bin/ArcView/firexaq?MERGE=1#TOGA.DC8_MRG/](https://www-air.larc.nasa.gov/cgi-bin/ArcView/firexaq?MERGE=1#TOGA.DC8_MRG/). More details on the
296 NASA online merging tool can be found in [https://www-
297 air.larc.nasa.gov/missions/etc/onlinemergedoc.pdf](https://www-air.larc.nasa.gov/missions/etc/onlinemergedoc.pdf).



309 **Figure 3.** Example time series of CAMS and TOGA-TOF HCHO and LGR CO observations during FIREX-AQ
 310 transverse (i.e., perpendicular to the transport of the fire plume) plume transects with relatively (a) high and (b) low
 311 variability in the observed 1 Hz HCHO. The pink shading shows when the DC-8 was inside the smoke plume based
 312 on the relative changes in CO. Vertical error bars on the TOGA-TOF HCHO data points show the measurement
 313 uncertainty ($\pm 35\%$), centered on the TOGA-TOF sampling time midpoint, and the horizontal error bars span the
 314 start to stop times of each TOGA-TOF sample. For each of the start-to-midpoint and midpoint-to-stop times of the
 315 TOGA-TOF sampling periods, 50% of the total sample volume occurred. These times (start/midpoint/stop) were
 316 used for deriving weighted averages of the 1 Hz HCHO and CO data.

312 The second averaging method is the volume-weighted averaging method based on the
 313 volume of air sampled. A post-mission laboratory analysis of the normalized volume vs. sample
 314 collection duration demonstrated that the flow rate through the sample trap was not constant during
 315 the sampling time period, despite the mass flow controller (MFC) being set to a fixed value.
 316 Specifically, relative flow rates were higher during the initial seconds of sampling before

317 decreasing to the MFC setpoint. Through a series of experiments, we quantified the total sample
318 volume as a function of collection time and derived an empirical relationship to determine the
319 sample-volume midpoint, defined as the time at which 50% of the total collected volume had been
320 accumulated during the sampling cycle (**Fig. 3**). Midpoints were calculated for each sample using
321 this method. The midpoints determined through this method are reported in the TOGA-TOF
322 FIREX-AQ data files archived at the NASA Atmospheric Science Data Center (ASDC)
323 (https://doi.org/10.5067/ASDC/FIREXAQ_TraceGas_AircraftInSitu_DC8_Data_1). The
324 arithmetic mean of the 1 Hz data is thus calculated for the time periods between the start to
325 midpoint (T_1) and midpoint to stop times (T_2), and these two means are averaged to weight the
326 merge value on the sampled volume.

327 When sampling near strong VOC sources (e.g., wildfires or urban emissions), air masses
328 often have significant heterogeneity of HCHO levels from varying direct emissions and
329 photochemical production from rapidly changing VOC precursors. In those cases, using the non-
330 weighted averages of 1 Hz measurements can lead to significant biases when they are directly
331 compared to the TOGA-TOF data. Therefore, using volume-weighted averages of higher-
332 frequency data that factor in the true sample volume midpoints for comparisons with TOGA-TOF
333 data allows a more accurate comparison of the sampled air masses.

334 Using the above methodology, volume-weighted averages of LGR and DACOM 1 Hz CO
335 data are carried out, during the times when TOGA was sampling. In our study, we use the volume-
336 weighted averaged LGR CO data when it is available, and for times when it is not available, we
337 use the volume-weighted averaged DACOM CO data multiplied by 0.978 to account for the
338 relative differences in the two CO datasets. We normalized to the LGR CO data as it is a little
339 lower to be conservative in our analyses, although the two CO datasets are in excellent agreement

340 (Fig. S4) and within the combined measurement uncertainties (LGR = 2 ppb + 2% and DACOM
341 = ± 2%).

342 2.4 Formaldehyde Normalized Excess Mixing Ratios

343 We compare HCHO normalized excess mixing ratios (NEMRs) for Western U.S. wildland
344 fires calculated using HCHO reported by TOGA-TOF, ISAF, and CAMS. There are several ways
345 to derive NEMRs (Yokelson et al., 2013a) and here we used the slope method, which involves
346 determining the slope of HCHO in ppt against CO in ppb for a single plume transect and the
347 background air outside the plume. CO was used as the reference species to normalize for dilution
348 as it is a conserved biomass burning tracer in wildfire plumes (Müller et al., 2016; Selimovic et
349 al., 2019; Yokelson et al., 2013a). For the NEMRs from CAMS and ISAF, we use the 1 Hz HCHO
350 and CO data for the entire plume transect. For the TOGA-TOF NEMRs, we use the discrete HCHO
351 sampling points, which range between two and ten samples for each plume transect — including
352 a background sample and plotted these against the volume-weighted average (described in Sect.
353 2.3) TOGA merged CO for corresponding sampling periods.

354 The NEMR analysis was carried out for the plumes selected based on the following criteria:
355 a) plume transects perpendicular to the fire source, b) availability of at least one full TOGA-TOF
356 sampling cycle within the plume, c) stable background HCHO prior to and after the plume transect,
357 and d) availability of both CAMS and ISAF 1 Hz HCHO data sets and CO for both background
358 and in-plume sampling. Fire plumes were identified via plume flags using above background CO
359 in addition to visual inspection of the data (FIRE_FLAG_TABULAR_DATA.dat, [https://www-
360 air.larc.nasa.gov/cgi-bin/ArcView/firexaq#SCHWARZ.JOSHUA/](https://www-air.larc.nasa.gov/cgi-bin/ArcView/firexaq#SCHWARZ.JOSHUA/)). Based on these criteria,
361 NEMRs were calculated for a total of 86 plume transects from 7 fire sources in the Western U.S.
362 and these are summarized in the supporting information (Table S1).

363 3. Results and Discussion

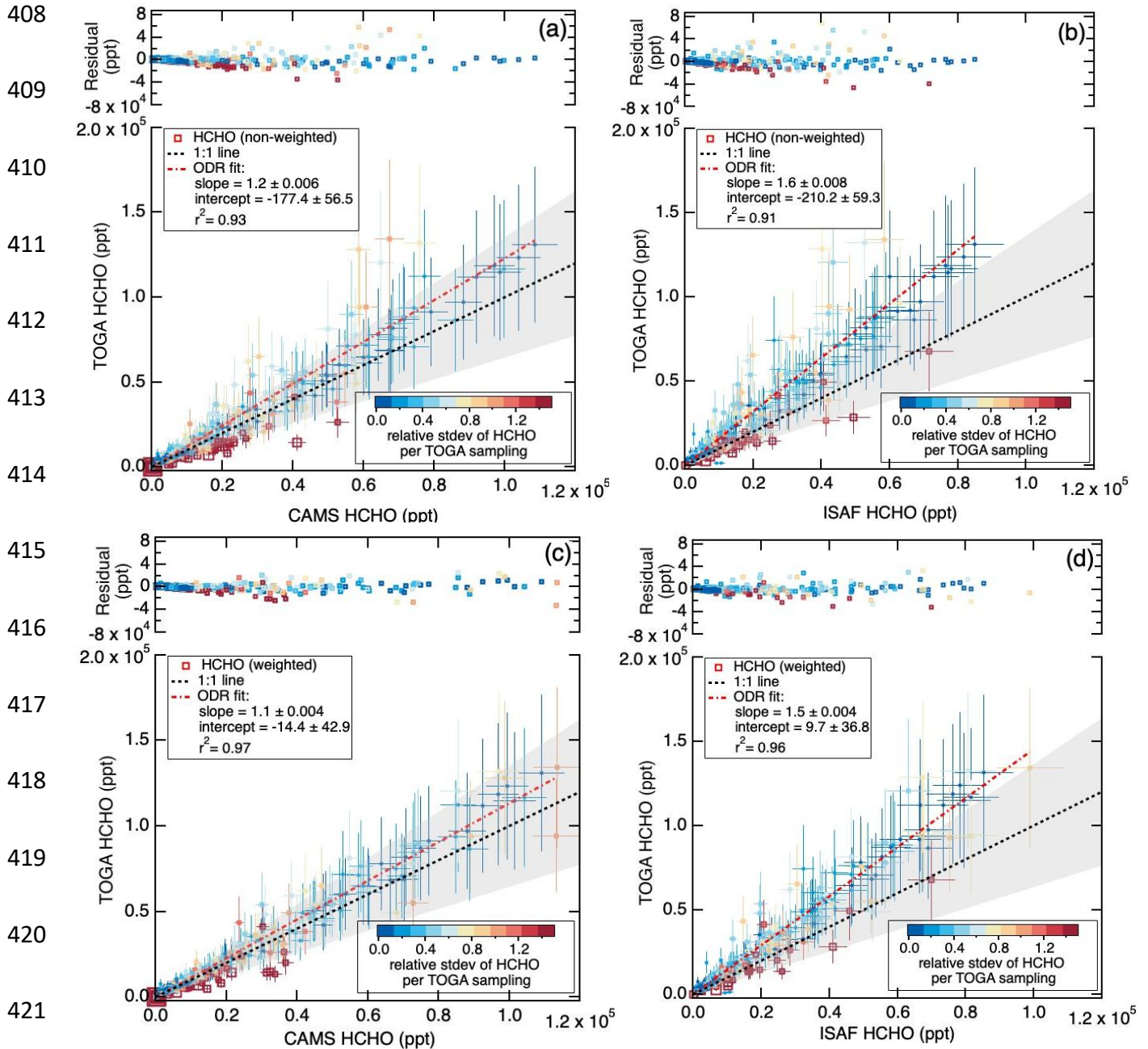
364 3.1 Comparison of HCHO between different instrumental techniques

365 Direct intercomparisons of measured HCHO between TOGA-TOF and the 1 Hz
366 instruments were carried out for data collected during all flights with available HCHO observations
367 onboard the DC-8. This included 13 flights sampling Western U.S. fires, 7 flights in the
368 Southeastern U.S., and 2 transit flights (**Fig. S1**). The dataset allowed comparisons of the widest
369 possible range of HCHO observed in the troposphere (below 100 ppt to ~ 100 ppb). **Figure 4**
370 shows correlations between TOGA-TOF, CAMS, and ISAF HCHO, averaged during the
371 corresponding TOGA-TOF sampling periods. Comparisons with normal averaged 1 Hz HCHO
372 (**Fig. 4a** and **b**) and volume-weighted averaged 1 Hz HCHO (**Fig. 4c** and **b**), described in **Sect.**
373 **2.3**, are both shown. The data points are colored according to the relative standard deviation of
374 HCHO, calculated as the standard deviation of 1 Hz HCHO data during the TOGA-TOF sampling
375 period divided by the HCHO measured by TOGA-TOF. This relative standard deviation represents
376 the variability of the 1 Hz HCHO data during the TOGA-TOF sampling period.

377 Both non-weighted (**Fig. 4a**) and volume-weighted (**Fig. 4c**) averages showed that HCHO
378 measured by CAMS was within the 35% measurement uncertainty of the TOGA-TOF while the
379 comparisons with ISAF (**Fig. 4b** and **d**) were outside of the range of measurement uncertainties.
380 When using the non-weighted averaging method (**Fig. 4a** and **b**), the bivariate fitting (orthogonal
381 distance regression, ODR) of the TOGA-TOF and the 1 Hz HCHO gave a slope of 1.2 for CAMS
382 (y -intercept -177.4 ppt, **Fig. 4a**) and 1.6 for ISAF (y -intercept -210.2 ppt, **Fig. 4b**). The y -axis
383 residuals from the ODR fit were largest for data points with higher relative standard deviations
384 (i.e., > 0.7) of HCHO. For the non-weighted averages, the absolute values of the y -axis residuals
385 ranged from 0.6 ppt to 59.0 ppb for CAMS (mean \pm standard deviation = 1.0 ± 3.0 ppb) and 0.1

386 ppt to 68.1 ppb for ISAF (0.9 ± 3.2 ppb). To differentiate the dependence on HCHO variations
387 during each TOGA-TOF sampling, a relative standard deviation of 0.7 was used as a cut-off
388 threshold. For data points with relative standard deviations > 0.7 , residuals ranged from 2.6 ppt to
389 59.0 ppb (2.7 ± 6.8 ppb) for CAMS and 52.1 ppt to 46.8 ppb (3.7 ± 7.8 ppb) for ISAF. In
390 comparison, for less heterogeneous air masses (i.e., TOGA-merge 1 Hz HCHO with relative
391 standard deviations < 0.7), residuals ranged from 0.6 ppt to 44.1 ppb (0.7 ± 1.8 ppb) for CAMS
392 and from 0.1 ppt to 68.1 ppb (0.6 ± 2.5 ppb) for ISAF. Therefore, the data points with less variable
393 HCHO (i.e., HCHO relative standard deviation < 0.7) showed on average a factor of 4–6 lower
394 residual values, showing that the increased heterogeneity of HCHO in the sampled air masses lead
395 to poorer correlations between TOGA-TOF and the 1 Hz HCHO instruments.

396 For more accurate comparisons, the CAMS and ISAF HCHO data averages were weighted
397 by the TOGA-TOF sampling volume, factoring in the 50% volume sampling midpoint. Weighted
398 averages of the CAMS and ISAF measured HCHO improved the correlations with TOGA-TOF to
399 give an r^2 of 0.97 (compared to 0.93 for non-weighted averages) for CAMS and 0.96 (compared
400 to 0.91 for non-weighted averages) for ISAF (**Fig. 4c** and **d**). The residuals of the volume-weighted
401 averages were 30% lower for CAMS (range 0.2 ppt to 32.2 ppb and average 0.7 ± 2.0 ppb) and
402 33% lower for ISAF (range 0.03 ppt to 32 ppb and average 0.6 ± 2.0 ppb). Biases at low mixing
403 ratios significantly improved when using volume-weighted averages with absolute values of the
404 y-intercepts much closer to 0 than the non-weighted averages, at -14.4 ± 42.9 ppt for CAMS
405 (versus -177.4 ± 56.4 ppt) and 9.7 ± 36.8 ppt for ISAF (versus -210.2 ± 59.3 ppt). Comparisons
406 between TOGA-TOF and the UiO PTR-ToF-MS (**Fig. S5**), showed similar results with better
407 agreement when using volume-weighted averages.



422 **Figure 4.** Correlation of measured HCHO (a,c) between TOGA-TOF and CAMS and (b,d) between TOGA-TOF
 423 and ISAF. CAMS and ISAF 1 Hz HCHO data were averaged over the TOGA-TOF sampling start and stop times
 424 using the (a,b) non-weighted and (c,d) volume-weighted averaging methods. Measurement uncertainties of each
 measurement technique are shown as error bars. The grey shading around the 1:1 line is the HCHO measurement
 uncertainties of the two corresponding instruments added in quadrature. Each data point is colored by the relative
 standard deviation of the 1 Hz HCHO data (standard deviation of 1 Hz averaged HCHO divided by TOGA-TOF
 HCHO) during the TOGA-TOF sampling period.

425 The discrepancies of HCHO between different measurement techniques during FIREX-
 426 AQ are similar to those shown by Liao et al. (2021), in which a comparison of CAMS vs. ISAF

427 HCHO data had a slope of 1.27. In our study, a comparison of the volume-weighted TOGA-merge
428 CAMS vs. ISAF HCHO from FIREX-AQ has a slope of 1.28, similar to Liao et al. (2021). The
429 minor difference in the slopes is most likely due to the different periods of data selected for analysis
430 as we only included CAMS and ISAF data during TOGA sampling in our analysis. Liao et al.
431 (2021) suggested that the main discrepancies between CAMS and ISAF were driven by the
432 uncertainties in the absolute calibration of the HCHO standards. As mentioned in **Sect. 2.2**, each
433 HCHO measurement technique has been carefully calibrated with HCHO standards. The HCHO
434 calibration standards used by each group were gravimetrically prepared and the absolute
435 concentrations were quantified using optical absorption techniques. As the main purpose of this
436 paper is to validate the TOGA-TOF as a unique technique for measuring HCHO, we will not
437 further explore the reasons for the discrepancies between the instruments.

438 **3.2 Comparison of HCHO NEMR between different instrumental techniques**

439 Normalized excess mixing ratios (NEMR) are used to derive the enhancement of a trace
440 gas within a smoke plume (Yokelson et al., 2013a). **Figure 5** shows HCHO NEMRs derived from
441 Western U.S. fires using the TOGA-TOF, CAMS, and ISAF instrumental techniques with respect
442 to smoke plume physical age ([https://www-air.larc.nasa.gov/cgi-](https://www-air.larc.nasa.gov/cgi-bin/ArcView/firexaq#SCHWARZ.JOSHUA/)
443 [bin/ArcView/firexaq#SCHWARZ.JOSHUA/](https://www-air.larc.nasa.gov/cgi-bin/ArcView/firexaq#SCHWARZ.JOSHUA/)). Based on the instrument uncertainties of both CO
444 and the individual HCHO measurements, the uncertainties of the NEMRs are 35%, 6%, and 10%
445 for TOGA-TOF, CAMS, and ISAF, respectively. The HCHO NEMRs observed during FIREX-
446 AQ from TOGA-TOF were in the range of 12 to 30 ppt ppb⁻¹ CO. These NEMRs were within the
447 wide range observed from previous wildfire biomass burning studies from 2.2 to 46 ppt ppb⁻¹ CO
448 (e.g., Hornbrook et al., 2011 and references therein) and during prescribed burning events in the
449 Southeastern U.S. during FIREX-AQ that ranged from 16 to 29 ppt ppb⁻¹ CO (Travis et al., 2023).

450 The quadratic polynomial fitting of HCHO NEMRs from all three instruments showed that the
451 NEMRs increased up to ~ 5 h of plume physical age then declined. The age at which the HCHO
452 NEMRs from each instrument begin to decrease (i.e., the change in the NEMRs with respect to
453 plume age = 0) are all within 20 minutes of 5 h. This trend with respect to plume physical age
454 agrees with what Liao et al. (2021) reported for HCHO NEMRs measured by ISAF from individual
455 fire sources and is driven by the balance of production of HCHO, mainly from VOC oxidation by
456 OH, and loss of HCHO, mainly from photolysis, throughout the aging of plume (Liao et al., 2021).

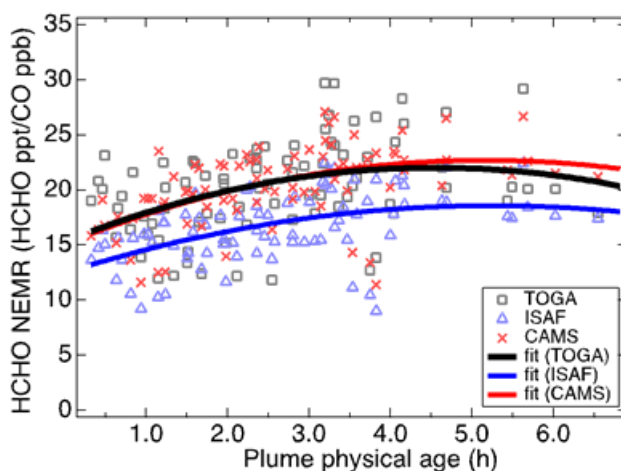
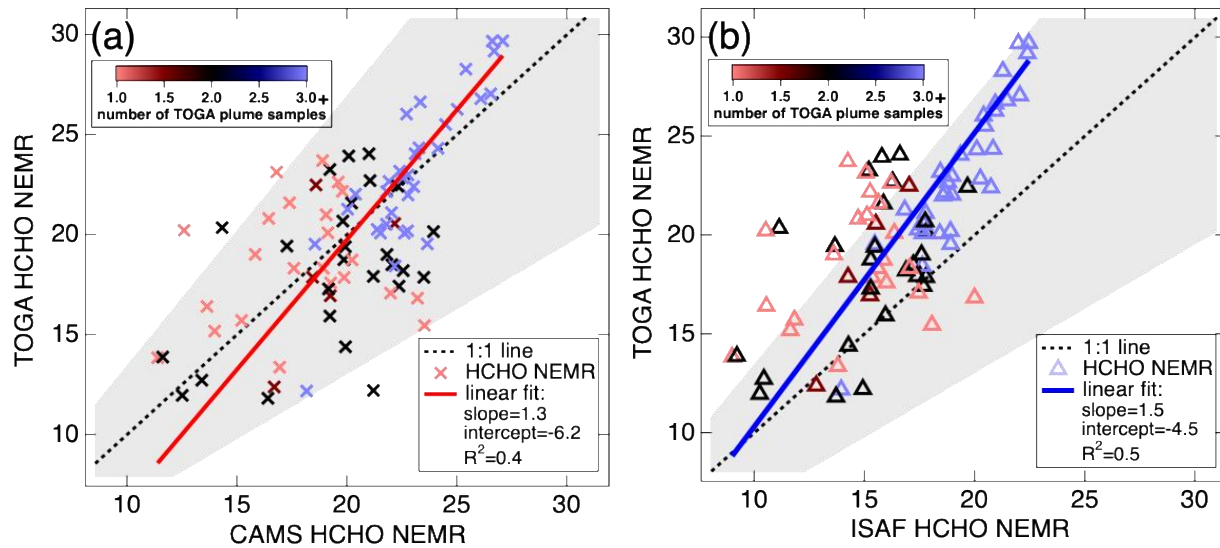


Figure 5. HCHO NEMRs of Western U.S. fires with respect to plume physical age during FIREX-AQ. Quadratic polynomial fittings of NEMRs derived using TOGA-TOF, CAMS, and ISAF HCHO data are shown.

457

458 While the trend with respect to plume physical age was similar using HCHO from the
459 different instruments, systematic differences were observed. To further investigate the possibility
460 of systematic differences of the NEMRs derived from the three HCHO instruments, paired *t*-tests
461 were used as a statistical method to compare two different analytical methods (Yen et al., 2020).
462 Here we carried out a paired *t*-test on the HCHO NEMRs calculated from the HCHO
463 measurements from the TOGA-TOF against CAMS and ISAF data. The sample number of the
464 analysis was equivalent to the number of selected transects for NEMR analysis (86 pairs). The null

465 hypothesis was whether the HCHO NEMRs from a pair of instruments are the same ($H_0: \mu_1 = \mu_2$),
 466 where the absolute value of the t -value (t) higher than the critical t (T_c) rejects the hypothesis.
 467 Between the TOGA-TOF and CAMS NEMRs, the analysis yielded $t = -0.42$, $T_c = 1.99$, and p -
 468 value = 0.68, retaining the hypothesis (at the $p = 0.05$ (5%) level). Between the TOGA-TOF and
 469 ISAF NEMRs, the analysis yielded $t = 11.53$, $T_c = 1.99$, and p -value = 4.68×10^{-19} , rejecting the
 470 hypothesis. Between the CAMS and ISAF NEMRs, the analysis yielded $t = 29.84$, $T_c = 1.99$, and
 471 p -value = 8.21×10^{-47} , again rejecting the hypothesis. Therefore, both TOGA-TOF and CAMS
 472 derived NEMRs had a systematic difference from the ISAF derived NEMRs with a significance at
 473 the $p = 0.05$ (5%) level.



474 **Figure 6.** Correlation plots between (a) TOGA-TOF and CAMS and (b) TOGA-TOF and ISAF derived HCHO
 NEMRs. Orthogonal distance regression fittings of the correlations are shown. The grey shading around the
 dashed 1:1 line shows the measurement uncertainty (35%) of TOGA-TOF HCHO.

475 As mentioned in **Sect. 3.1**, systematic biases exist between the HCHO measurement
 476 techniques which are likely due to calibration standard differences. As illustrated in **Fig. 5**, these
 477 biases are similarly present in the derived HCHO NEMRs. Nonetheless, comparisons between the

478 HCHO NEMRs derived from TOGA-TOF and CAMS (**Fig. 6a**) and TOGA-TOF and ISAF (**Fig.**
479 **6b**) indicate that the correlations are within the uncertainty bounds of the NEMR.

480 3.3 Discussion on TOGA-TOF HCHO NEMRs and their applicability

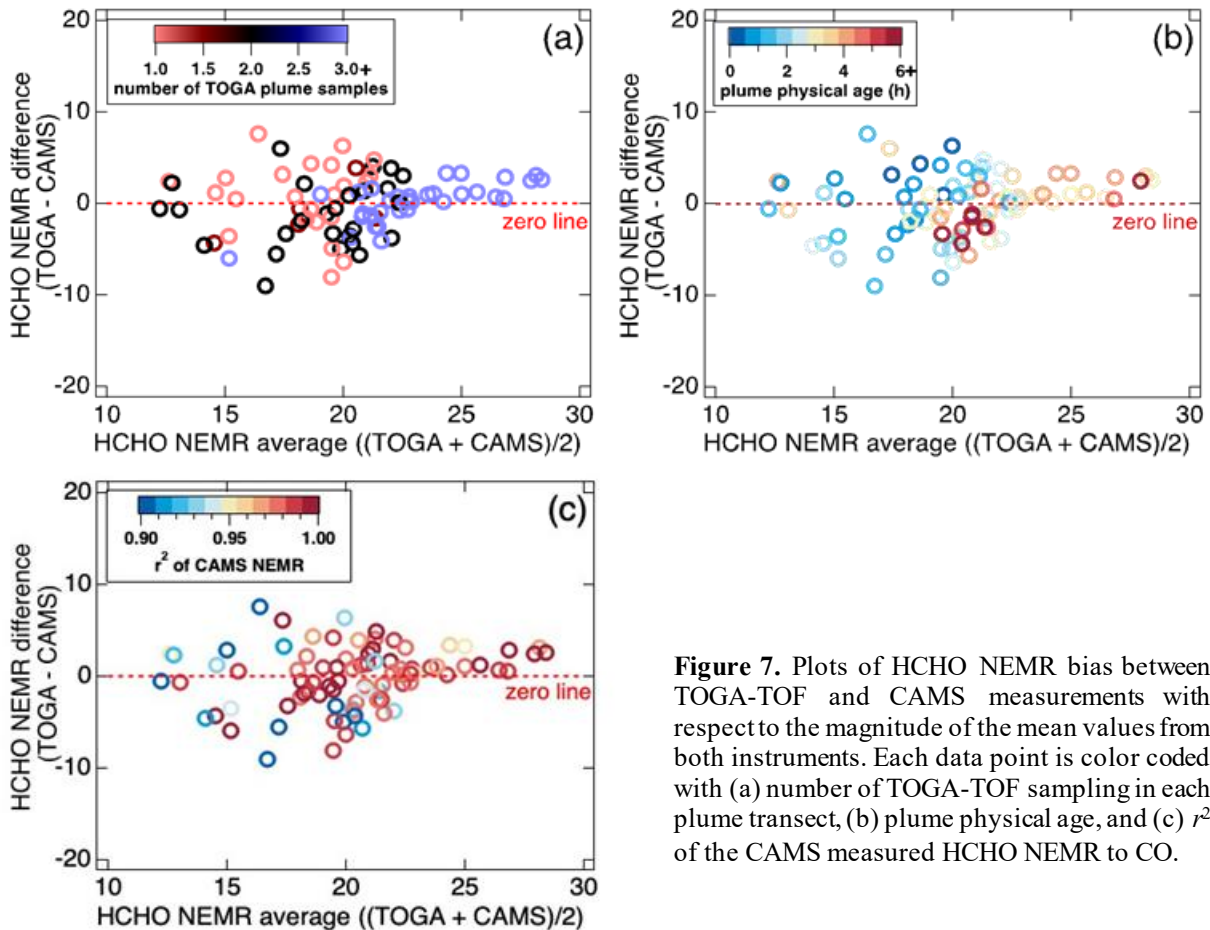


Figure 7. Plots of HCHO NEMR bias between TOGA-TOF and CAMS measurements with respect to the magnitude of the mean values from both instruments. Each data point is color coded with (a) number of TOGA-TOF sampling in each plume transect, (b) plume physical age, and (c) r^2 of the CAMS measured HCHO NEMR to CO.

481

482 The applicability of TOGA-TOF-derived HCHO NEMRs is further explored for different
483 smoke plume conditions. **Figure 7** shows the difference of HCHO NEMRs between TOGA-TOF
484 and CAMS with respect to the averages from the two methods. Similar to **Fig. 5**, **Fig. 7b** shows
485 that the averaged HCHO NEMRs increase with plume physical age. With greater plume physical
486 age, the bias between TOGA-TOF driven HCHO NEMRs to CAMS becomes lower (**Fig. 7b**) and
487 this lower bias corresponds to more TOGA-TOF samples at each plume transect (**Fig. 7a**). With

488 increasing physical age, the smoke plumes studied were generally more dispersed and spatially
489 wider, therefore allowing more TOGA-TOF samples per transect.

490 The reduced bias between the two instrumental techniques with plume physical age can be
491 explained from the reduced plume heterogeneity with aging, as indicated by the improved r^2 of
492 each plume sampling with plume physical age (**Fig. 7c**). Modeling studies of fire plumes
493 (Trentmann et al., 2003a, 2003b; Wang et al., 2021) have shown significant spatial heterogeneity,
494 both vertically and horizontally, of fire plumes with different levels of oxidants (i.e., OH, O₃, NO₃).
495 This plume heterogeneity has also been observed in power plant plumes (Brock et al., 2002). As
496 discussed in Wang et al. (2021), at the early stage of a fire plume, OH levels are severely dampened
497 at the core of the plume, mainly due to attenuation of solar radiation from aerosols, but enhanced
498 at the edges. With aging of the plume, OH levels become more homogeneous (Wang et al., 2021).
499 Liao et al. (2021) showed that HCHO levels observed in fire plumes as they age during FIREX-AQ
500 were primary emissions coupled with subsequent photolytic loss and secondary production,
501 primarily from the oxidation of VOCs with OH. Therefore, with plume physical age, as oxidant
502 levels and photochemistry become more homogeneous, we can expect HCHO NEMRs to be more
503 constant throughout a transect perpendicular to the direction of the smoke transport. As such, the
504 r^2 value of the slope of CAMS HCHO and CO, from which the NEMRs were calculated, is closer
505 to 1 for larger NEMRs (and larger plume physical ages) (**Fig. 7c**). These results demonstrate that
506 the TOGA-TOF data are sufficient for assessing HCHO NEMRs in plumes with physical ages >
507 2–3 h and may result in a small bias in calculated NEMRs for younger, narrower plumes in
508 comparison to those derived using fast HCHO measurement techniques.

509 4. Conclusions

510 In this study, we demonstrated the TOGA-TOF as a fast response (< 2 min) GC/MS
511 instrument that can measure HCHO along with > 100 additional C_1 to C_{10} VOCs. During FIREX-
512 AQ, the TOGA-TOF measured a wide range of HCHO mixing ratios with an LLOD of 20 ppt
513 aboard the NASA DC-8. To our knowledge, this technique has not been previously reported for
514 any ambient measurements. The discrete TOGA-TOF HCHO measurements during FIREX-AQ
515 are comparable to the averaged non-discrete 1 Hz HCHO data measured by the CAMS and UiO
516 PTR-ToF-MS instruments while the discrepancies are beyond the combined instrumental
517 uncertainties for the TOGA-TOF and ISAF instruments. The discrepancies between HCHO
518 instrumental techniques during FIREX-AQ are currently believed to be due to differences in
519 calibration standards as quantified by different optical techniques. Using volume-weighted TOGA
520 merge averages of HCHO measured by CAMS, ISAF, and UiO PTR-ToF-MS improved the
521 comparison agreement with TOGA-TOF HCHO observations. Therefore, we strongly encourage
522 the community to use the reported Time_Start, Time_Mid, and Time_Stop values in the TOGA-
523 TOF archived data files to determine volume-weighted averages of concurrent measurements.

524 TOGA-TOF derived NEMRs showed similar trends with plume physical age to what has
525 been reported from ISAF (Liao et al., 2021), with NEMRs increasing with plume ages up to ~ 5 h.
526 Due to its nature of taking discrete subsamples of air masses, as shown in Fig. 3a, highly
527 heterogeneous features may not be captured by the TOGA-TOF system in comparison to fast
528 HCHO measurement techniques, especially in chemically dynamic environments near strong
529 sources with highly heterogeneous emissions and rapidly varying HCHO production and loss often
530 detected during airborne measurements. Nevertheless, the technique used in the TOGA-TOF offers
531 a powerful HCHO-measuring method to the scientific community, with the capability to
532 simultaneously measure > 100 other C_1 - C_{10} VOC species. The wide range of gas-phase VOCs

533 measured with the TOGA-TOF can be used to provide significant insights into understanding the
534 photochemical state of air masses in rural to polluted environments. In addition, the time resolution
535 and inherent averaging is well-suited for verifying satellite observations, three-dimensional
536 modeling, and to compare with other instrumental techniques.

537 **Data availability.** All data used in this paper are available at [https://www-](https://www-air.larc.nasa.gov/missions/firex-aq/index.html)
538 [air.larc.nasa.gov/missions/firex-aq/index.html](https://www-air.larc.nasa.gov/missions/firex-aq/index.html)

539
540 **Author contributions.** ECA conceptualized the project. DJ performed the data analyses and wrote
541 the manuscript with input from ECA and RSH, RSH, AJH, GD, HSH, JPD, AF, DR, JW, PW,
542 TFH, GMW, JSC, JP, AW, TM, JBN, FP, LT, JD, CW, JS, ECA performed the measurements,
543 CDH analyzed the fire smoke ages, AS and EG contributed to the smoke and fuel identification
544 data product, JHC was the project PI and provided guidance on the analyses.

545
546 **Competing Interests.** The authors declare they have no conflict of interest. Some authors are
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548
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