1	Emissions of volatile organic compounds (VOCs) from concentrated
2	animal feeding operations (CAFOs): chemical compositions and
3	separation of sources
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18 Abstract

19 Concentrated animal feeding operations (CAFOs) emit a large number of volatile organic 20 compounds (VOCs) to the atmosphere. In this study, we conducted mobile laboratory 21 measurements of VOCs, methane (CH_4) and ammonia (NH_3) downwind of dairy cattle, 22 beef cattle, sheep and chicken CAFO facilities in northeastern Colorado using a 23 hydronium ion time-of-flight chemical-ionization mass spectrometer (H_3O^+ ToF-CIMS) 24 that can detect numerous VOCs. Regional measurements of CAFO emissions in 25 northeastern Colorado were also performed using the NOAA WP-3D aircraft during the 26 Shale Oil and Natural Gas Nexus (SONGNEX) campaign. Alcohols and carboxylic acids 27 dominate VOC concentrations and the reactivity of the VOCs with hydroxyl (OH) 28 radicals. Sulfur-containing and phenolic species provide the largest contributions to the 29 odor activity values and the nitrate radical (NO_3) reactivity of VOC emissions, 30 respectively. VOC compositions determined from mobile laboratory and aircraft 31 measurements generally agree well with each other. The high time-resolution mobile 32 measurements allow the separation of the sources of VOCs from different parts of the 33 operations occurring within the facilities. We show that the emissions of ethanol are 34 primarily associated with feed storage and handling. Based on mobile laboratory 35 measurements, we apply a multivariate regression analysis using NH_3 and ethanol as 36 tracers to determine the relative importance of animal-related emissions (animal 37 exhalation and waste) and feed-related emissions (feed storage and handling) for different 38 VOC species. Feed storage and handling contribute significantly to emissions of alcohols, 39 carbonyls, carboxylic acids and sulfur-containing species. Emissions of phenolic species 40 and nitrogen-containing species are predominantly associated with animals and their 41 waste.

42 **1. Introduction**

43 Concentrated animal feeding operations (CAFOs) emit many volatile organic 44 compounds (VOCs) into the atmosphere, including carboxylic acids, alcohols, carbonyls, 45 phenolic compounds, sulfur- and nitrogen-containing compounds (Hobbs et al., 46 2004; Filipy et al., 2006; Sun et al., 2008; Ni et al., 2012). These VOCs can contribute to 47 the formation of ozone (Howard et al., 2010a;Howard et al., 2010b;Gentner et al., 2014) 48 and fine particles (Sintermann et al., 2014; Perraud et al., 2015), both affecting regional 49 air quality. Many VOCs from CAFOs are also responsible for the unpleasant odor 50 problems nearby or downwind of these facilities (McGinn et al., 2003;Rabaud et al., 51 2003;Parker et al., 2010;Woodbury et al., 2015). Some VOCs (e.g. phenolic species) (US 52 EPA, 2016) from CAFOs are harmful to human health. 53 There are a large number of potential VOC sources inside a CAFO, potentially 54 including animal exhalation, animal waste in animal pens, flushing lanes, lagoons, silage 55 storage piles and silos, and feed mixtures in feed lanes and bunks (Alanis et al., 56 2008; Chung et al., 2010). Early studies mainly focused on VOC emissions from animal 57 waste (e.g. slurry and manure) under laboratory conditions (Hobbs et al., 1997, 58 1998;Hobbs et al., 2004). Ngwabie et al. (2007;2008) reported that VOC concentrations 59 in dairy, sheep and pig CAFOs were highest during animal waste removal and feeding, 60 indicating that large emissions were related to these activities. Recent studies found that 61 VOC concentrations in dairy farms were significantly higher near silage and piles of 62 animal feed (i.e. total mixed rations) than near other places (animal pens, lagoons and 63 flush lanes), suggesting that feed-related sources dominate VOC emissions (Alanis et al., 64 2008; Chung et al., 2010). Enhancements of some VOCs (e.g. acetone) in animal sheds 65 are also related to animal exhalation (Shaw et al., 2007;Ngwabie et al., 2008;Sintermann 66 et al., 2014). However, the contributions of different sources to individual VOC 67 emissions from a facility are not accurately known (Ngwabie et al., 2008). This poor 68 understanding of VOC sources hinders the development of management practices that 69 reduce VOC emissions in animal feeding facilities (Ngwabie et al., 2007). Thus, a 70 comprehensive characterization of VOC sources and their relative importance within a 71 CAFO is needed.

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Many studies of VOCs from animal feeding operations have been conducted with

73 offline analytical methods (Filipy et al., 2006; Alanis et al., 2008; Sun et al., 2008; Chung 74 et al., 2010; Ni et al., 2012). VOCs were collected on filters, or in canisters and cartridges 75 and were quantified in the laboratory using various methods (see reviews in Ni et al., 2012). These offline methods are labor-intensive, which limits the number of VOC 76 77 samples. Online fast measurement techniques (mainly proton-transfer reaction mass 78 spectrometers, PTR-MS) allow for more detailed investigation of CAFO facilities (Shaw 79 et al., 2007;Ngwabie et al., 2008;Sintermann et al., 2014). The previous online 80 measurements usually used a single stationary sampling inlet either inside a stall or at a 81 fence line, which does not provide spatial distribution information for VOCs in the 82 facilities.

In this study, we deployed a high time-resolution instrument on board a mobile laboratory driven on public roads and a NOAA WP-3D research aircraft to measure VOCs downwind of CAFO facilities. We will use this dataset to characterize chemical compositions of VOC emissions and explore different sources within the facilities that contribute to VOC emissions.

88 2. Experiments

89 Mobile laboratory measurements were conducted near Greeley in northeastern 90 Colorado, US. Six different CAFOs were studied, including two dairy farms, two beef 91 feed yards, one sheep feed yard and one egg-laying chicken farm (Table S1). Among the 92 six CAFOs, emissions of NH_3 , N_2O and CH_4 in four facilities (the two dairy farms, one 93 beef cattle feed yard and the sheep feed yard) have been measured previously using an 94 instrumented van (Eilerman et al., 2016). We added a new VOC instrument to the 95 payload, and performed mobile measurements in winter time (February, 2016) for the six 96 CAFOs by sampling at their downwind flanks 1-2 times for each facility. Duplicated 97 measurements at the same facilities agreed well.

98 VOCs were measured using a hydronium ion time-of-flight chemical-ionization 99 mass spectrometer (H_3O^+ ToF-CIMS) instrument on the mobile laboratory. Here, we 100 provide a brief description of the instrument (see details in Yuan et al., 2016). VOCs are 101 ionized by H_3O^+ ions in a drift tube, similar to a PTR-MS (de Gouw and Warneke, 2007). 102 The protonated product ions are detected using a high-resolution time-of-flight (ToF) 103 analyzer (Tofwerk AG) (m/ Δ m=4000-6000). A number of VOC species were calibrated

104 using either gravimetrically prepared gas cylinders or permeation tubes (see details in

105 Yuan et al., 2016). VOC background signals in the instrument were determined by

106 passing ambient air through a catalytic converter. The detection limits are compound-

107 dependent and range between 10-100 ppt for most VOC species at a time resolution of 1

108 s. Besides VOCs, two inorganic species, NH_3 and H_2S , were measured at m/z 18.034

109 (NH₄⁺) and m/z 34.995 (H₃S⁺) using the H₃O⁺ ToF-CIMS, respectively (Li et al.,

110 2014;Müller et al., 2014).

111 In addition to the H_3O^+ ToF-CIMS, a cavity ringdown spectrometer (CRDS) 112 instrument (Picarro G1301m) measuring methane (CH_4) and carbon dioxide (CO_2) and an 113 off-axis integrated cavity output spectrometer (OA-ICOS, Los Gatos Research) 114 measuring nitrous oxide (N_2O) along with carbon monoxide (CO) were deployed during 115 the mobile laboratory measurements. Measurements of ambient temperature, relative 116 humidity, wind direction, wind speed and vehicle location were performed using 117 meteorological sensors (R.M. Young 85004 and AirMax 300WX) and a GPS compass 118 system (ComNav G2B). A summary of meteorological conditions during the mobile 119 laboratory measurements is shown in Table S2.

120 Measurements of agricultural plumes were also performed using the NOAA WP-3D 121 research aircraft in March-April, 2015 during the Shale Oil and Natural Gas Nexus 122 (SONGNEX) campaign. Data from three flights (March 28, March 29, April 13) over northeastern Colorado are used in this study. VOCs were measured using the same H_3O^+ 123 124 ToF-CIMS instrument as mobile measurements (Yuan et al., 2016). Another chemical 125 ionization mass spectrometer (CIMS) was used to detect NH₃ during SONGNEX (Nowak 126 et al., 2007). Due to background issues and lower concentrations, NH_3 signals were not 127 retrievable from H_3O^+ ToF-CIMS during the SONGNEX campaign.

3. Results and Discussions

129 **3.1.** Spatial distributions from mobile laboratory measurements

Figure 1 shows measured concentrations of NH₃, CH₄, CO₂, N₂O, ethanol (C₂H₅OH),
acetic acid (CH₃COOH) and acetone (CH₃COCH₃) around a beef feed yard (beef #1). The
concentrations of the seven species were enhanced and highly variable downwind of the

133 facility. Different time variations for the seven species were clearly observed. NH_3

134 concentrations peaked at around 13:46 local time (LT) and the peak location was

135 northwest of this facility, directly downwind of the animal pens. This is consistent with 136 fresh waste of animals (urine and feces) as the main source of NH₃ within a CAFO 137 facility (Hristov et al., 2011). CO₂ and CH₄ are emitted from animal respiration and eructation of the cattle (Shaw et al., 2007; Sintermann et al., 2014; Owen and Silver, 138 139 2015). CO₂ (R=0.77) and CH₄ (R=0.77) correlated well with NH₃ between 13:42-13:46 140 when NH_3 was high. These observations reflect the fact that animals and their fresh waste 141 may be largely co-located in the animal pens. But, waste cleaning time/practices in the 142 facility were unknown, owing to no access to the facility. Previous mobile and aircraft 143 measurements have also observed enhancements of NH₃ and CH₄ concentrations 144 downwind of animal pens in cattle feedlots (Miller et al., 2015;Hacker et al., 2016). The 145 time variations of two VOCs, acetic acid and acetone, followed reasonably well with both 146 NH₃ and CO₂, suggesting that animals and their waste contributed to the enhancements of 147 the two VOCs. Based on previous studies, the emissions from animal respiration should 148 dominate over waste for acetone, and vice versa for acetic acid (Ngwabie et al., 149 2008;Sintermann et al., 2014). The similar time variations of NH₃, CO₂, acetone and 150 acetic acids (and less clearly for CH₄) imply that the co-located emissions from animals 151 and their waste may not be separated based on the variations observed (over a short time 152 span of several minutes) while measuring along the downwind flanks.

153 A single, narrow high concentration spike (up to 1 ppm) of ethanol was observed 154 around 13:44 LT (Figure 1). The hotspot of ethanol was located downwind of a feed mill 155 at the west side of the feed yard, indicating that the feed mill and its related activities can 156 emit large amounts of ethanol. Distillers grains, a fermented by-product from ethanol 157 production, are commonly used as an ingredient of feed in beef cattle feed yards (Raabe, 158 2012) (although not known specifically here). Therefore, it is not surprising to observe 159 large emissions of ethanol in the feed mill area. High concentrations of many other VOC 160 species (e.g. acetic acid and acetone) were observed in the feed mill plume, whereas NH₃, 161 CH₄ and N₂O were not enhanced. Combustion sources, possibly due to equipment 162 operation in the feed mill area, are likely responsible for the enhancement of CO₂ and CO 163 at \sim 13:44 LT. VOC emissions from these combustion plumes are negligible (see details 164 in SI, Figure S1).

165 Measurements downwind of a dairy farm (dairy #1) are shown in Figure 2. The

highest concentrations of NH₃ and CO₂ were observed downwind of the animal pens, 166 167 similar to the beef feed yard shown in Figure 1. Interestingly, several high concentration 168 peaks of ethanol were observed along the drive track of the mobile laboratory. These peaks were in close proximity or downwind of the feed lanes (white lines on the satellite 169 170 image, Figure 2B). As shown for beef #1, feed mills can be an important source of 171 ethanol and other VOCs. Different from the usage of distillers grains in beef cattle feed 172 yards, silage is more commonly used as fodder for dairy cattle (Raabe, 2012). Previous studies showed that ethanol is the most abundant VOC species emitted from feed silage 173 174 (Hafner et al., 2013). It is expected that VOCs will continue evaporating from the feed 175 mixtures after the feed is delivered to the feed lanes. Time variations of acetic acid (and 176 acetone) correlated more closely with ethanol (R=0.72) than with NH₃ (R=-0.30) and CO₂ (R=-0.14), which differs from the beef feed yard. This suggests that the three VOCs were 177 178 mainly from emissions of feed lanes, rather than animals and their waste in this dairy 179 farm.

180 In addition to animals and their waste (referred to as animal+waste hereafter) and 181 feed storage and handling (referred to as feed storage+handling hereafter), we identified 182 another important VOC source from the other dairy farm studied (dairy #2, Figure S2). 183 High concentrations of ethanol, acetone, dimethyl sulfide (DMS, C_2H_6S) and CH_4 were 184 observed downwind of three milking parlors. Acetic acid was only moderately elevated, 185 whereas NH₃ was not enhanced. Compared to feed storage+handling, emission 186 compositions from the milking parlors are different. The emissions from milking parlors 187 might result from several sources, including animal exhalation and milking-related 188 activities. It is worth noting that we did not distinctly observe emissions from the milking 189 parlor in the dairy farm #1, which were potentially mixed with emissions from feed prior 190 to sampling.

191 The measurements downwind of three other CAFO sites (beef #2, the sheep feed 192 yard and the chicken house) are investigated in a similar way, as shown in Figure 1-2 193 (and Figure S2-S3). From this analysis, we identify three main VOC emission sources in 194 animal feeding facilities, namely animal+waste, feed storage+handling and milking 195 parlors. These measurements suggest that combustion sources are not important for VOC 196 emissions in these facilities.

197 **3.2. VOC chemical compositions of different CAFOs**

198 The enhancements of VOCs downwind of each CAFO were integrated to determine 199 the averaged VOC compositions for each facility (Figure 3). The measured VOC species 200 are divided into six different groups, namely carboxylic acids, alcohols, carbonyls, 201 phenolic species, nitrogen- and sulfur-containing species. The averages of the sum of 202 measured VOC concentrations downwind of the sites are in the range of 22-139 ppb, with 203 higher concentrations at dairy farm #1 and the two beef feed yards (wind speeds during 204 the measurements were similar except at the chicken house, Figure 3E and Table S1). As 205 demonstrated in Figure 3A, alcohols (55-87%, mole fractions) and carboxylic acids (4-206 32%) represent major classes of VOC from these CAFOs. Other VOC classes account for 207 8-21% of VOC concentrations in total.

208 As discussed in Section 1, VOC emissions from CAFOs can contribute to 209 unpleasant odor problems and ozone formation. We utilize odor activity value (OAV) 210 and the OH and NO₃ reactivity to evaluate the relative contribution of each VOC class to 211 the two environmental effects, respectively. The dimensionless OAV is estimated from 212 VOC concentrations divided by the species' single compound odor thresholds (SCOT) (OAV_i=C_i/SCOT_i) (Feilberg et al., 2010;Parker et al., 2010;Woodbury et al., 2015). The 213 214 reported SCOT values in the literature are highly variable and we use the geometric 215 means of literature values for each compound, as compiled in Parker et al. (2010). The 216 averaged total OAVs from all measured VOCs are in the range of 0.8-5.5 at different 217 sites (Figure 3B). Sulfur-containing species contribute the largest fractions (51-91%) to 218 total OAVs at different sites, followed by phenolic species (6-37%) and carboxylic acids 219 (3-11%). The relative contributions of different species to OAV agree well with previous estimates based on measurements from a beef feed yard (Woodbury et al., 2015). 220 221 OH reactivity (OHR) and NO₃ reactivity (NO3R) are determined as the products of 222 VOC concentrations and the respective reaction rate constants of VOCs with the two 223 oxidants (Atkinson et al., 2006) (OHR_i = $C_i \times k_{OH_i}$; NO3R_i = $C_i \times k_{NO3_i}$). The averaged OH reactivities range between $1-10 \text{ s}^{-1}$, which is comparable or lower than the typical OH 224 reactivity observed in urban areas (a few s⁻¹ to 50 s⁻¹) (Yang et al., 2016). Alcohols are the 225 226 largest contributors (40-75%) to OH reactivity at the sites, although the fractions from 227 carbonyls, phenolic and sulfur-containing species are also significant (Figure 3C). These

results are generally consistent with the finding that ethanol accounts for the majority of

- ozone formation potential of VOC emissions from a dairy farm (Howard et al., 2008).
- **230** The averaged NO₃ reactivities range from 0.02 to 0.26 s⁻¹, which are remarkably higher

than in urban areas (usually $<0.01 \text{ s}^{-1}$) (Tsai et al., 2014;Brown et al., 2016). In contrast to

the OH reactivity, phenolic species account for the largest fractions (66-90%) of the NO_3

reactivity for all of the sites, with remaining contributions primarily from sulfur- and

nitrogen-containing species (Figure 3D). We note that OAV, OH and NO₃ reactivity are

235 measured along the fence line and they decrease rapidly with downwind distance and

dilution (see example in section 3.4 for aircraft measurement results associated with a

factor of ~ 10 lower concentrations than those from mobile laboratory).

238 **3.3. Relative importance of different sources for VOC emissions**

239 As shown in section 3.1, ethanol was primarily emitted from feed storage+handling 240 (and milking parlors), whereas NH₃ and CO₂ were attributed to emissions from animals 241 and their waste. This suggests that these species can be used as tracers to separate the 242 emissions from sources. However, there are two issues that need to be considered: (1) 243 Emissions of animal exhalation and waste are largely co-located in the animal pens. As CO_2 is also emitted from combustion sources (see details in SI) and animal exhalation is 244 245 only important for a few species (e.g. acetone), NH₃ will be used as a tracer for the 246 emissions from animals and their waste. It is worth mentioning that long-term 247 measurements in CAFO facilities could permit separation of the two co-located sources 248 (see example in Sintermann et al., 2014). (2) There is some ethanol attributable to 249 animal+waste emissions that needs to be accounted for. Ethanol concentrations solely 250 from feed emissions ($[C_2H_5OH]_{Feed}$) can be calculated by subtracting the contribution of 251 ethanol by animal+waste from measured ethanol concentrations (see details in SI, Figure 252 S4).

After correcting ethanol for animal+waste emissions, the contributions of emissions from feed storage+handling and animal+waste to measured VOC enhancements at each individual site can be determined using multivariate linear fits to $[C_2H_5OH]_{Feed}$ and NH₃ concentration ($[NH_3]$).

257 $[VOC] = ER_{C_2H_5OH} \times [C_2H_5OH]_{Feed} + ER_{NH_3} \times [NH_3] + [bg]$ Eq. (1) 258 Here, [VOC] and [bg] are measured concentrations of the VOC species and the background concentration outside the CAFO plumes, respectively. $ER_{C_2H_5OH}$ and ER_{NH_3}

are the emission ratios of the VOC species relative to ethanol and NH_3 from the

emissions of feed storage+handling and animal+waste, respectively. Along with [bg], the emission ratios are determined from the multivariate linear fits.

Based on the fitted parameters from Eq. 1 (and Eq. S3 for dairy farm #2), the relative contributions of different sources to the enhancements of various VOC species can be calculated for the investigated sites (Figure 4). In general, large differences in fractional contributions to VOC enhancements exist among both different VOC species and different animal types. The main findings from Figure 4 are:

268 (1) Phenol, cresols, butanediones and many nitrogen-containing species are primarily269 associated with animal+waste emissions for the investigated sites.

270 (2) Both feed storage+handling and animal+waste account for significant fractions of
271 emissions of many oxygenated VOCs and sulfur-containing species.

272 (3) Based on the results from the dairy farm #2, emissions from milking parlors

273 contribute significantly to the enhancements of a limited number of VOC species,

including ethanol $(23\pm1\%)$, acetone $(35\pm3\%)$, acetaldehyde $(31\pm3\%)$, methanol $(18\pm3\%)$,

275 MEK (14±2%) and DMS (14±2%).

276 (4) Feed storage+handling plays important roles in the emissions of many VOC277 species from the chicken farm. Based on a news report on the facility, a manure belt

278 system is used to manage manure in this facility. The manure belt system catches the

279 excreta from chicken to transport manure to a separate location for storage. The chicken

houses with manure belt usually lead to substantially lower emissions (e.g. NH₃) from

animal waste (Wood et al., 2015). It is consistent with significantly lower NH₃

concentrations (0-175 ppb, Figure S3) at this site compared to ruminant feed yards

283 measured in this study (0-1000 ppb), although wind speed was 36%-60% higher during

284 measurements of the chicken house (7.5 m/s) than others (4.7-5.5 m/s). It is also possible

that emissions of NH_3 and VOCs were treated when in-house air was ventilated out

286 (Wang et al., 2010).

287 We further determine the contributions of emissions from feed storage+handling,

animal+waste and milking parlors to the total VOC concentrations (first columns in

Figure 4, also Figure S5). Feed storage+handling emissions account for 35-41%, 23-30%,

13% and 41% of the summed total VOC concentrations for the investigated dairy farms,
beef feed yards, the sheep feed yard and the chicken farm in this study, respectively. The
fractional contributions from the sources to odor activity value, OH reactivity and NO₃
reactivity are also calculated (Figure S5). The contributions from feed storage+handling
emissions to the three parameters are generally comparable or slightly smaller than those
contributions to the total VOC concentrations.

296 In addition to the information on relative contributions from different sources, the 297 multivariate fit analysis also provides the emissions ratios of VOCs to NH_3 for 298 animal+waste and emission ratios of VOCs to ethanol for both feed storage+handling and 299 milking parlors (see Table S3-S8). These emission ratios represent chemical 300 "fingerprints" of the emissions from various sources. The emission ratios are summed up 301 for the VOC classes and the fractions of each VOC class in different source emissions are 302 determined (Figure 5). Overall, VOC emissions from both feed storage+handling and 303 milking parlors are dominated by alcohols, whereas the contributions of carboxylic acids 304 and other VOC classes are significantly larger for animal+waste emissions. The VOC 305 compositional fractions shown in Figure 3A for each site are the weighted average of the 306 fractions for different sources in Figure 5.

We acknowledge that there are some limitations in separating different sources inside
each facility using measurements from the mobile laboratory, which may introduce some
uncertainties to the results.

310 (1) In this study, the relative fractions of different sources to VOC emissions are 311 determined based on snapshots of measurements when the mobile laboratory passed by 312 the CAFO sites. The relative fractions may change over time, and may be related to 313 operation activities within the facilities, such as feed mixing activities in feed mill area. 314 Nevertheless, some encouraging evidence was observed: the determined relative fractions 315 from different sources are reasonably similar between the two beef feed yards. The 316 agreements between the two dairy farms are not as good as for the two beef feed yards. 317 The above mentioned observations of the emissions from milking parlors and potential 318 differences in feed ingredients for dairy cattle, which are reflected by the discrepancies in 319 VOC compositions emitted from feed storage+handling (Figure 5A), could be the 320 reasons.

(2) VOCs from various sources in a CAFO site are mostly emitted at the surface.
VOCs were measured on the van at a single level near-ground (~3 m). However, the
plumes from CAFOs become deeper as they are transported downwind, and VOC
concentrations are vertically diluted by background air due to turbulent mixing. As
shown in Figure 1-2 (and Figure S2-S3), the feed mills and milking parlors at the sites
studied in this work are located nearby public roads, and their contributions may be
somewhat overestimated as a result.

328 3.4. Aircraft measurements

329 Time series of NH_3 and several VOCs inside two agricultural plumes in northeastern 330 Colorado measured on April 13, 2015 from the NOAA WP-3D during the SONGNEX 331 campaign are shown in Figure 6. Large enhancements of NH_3 were observed in the two 332 agricultural plumes, although the peak concentrations were a factor of ~10 lower than 333 those from mobile laboratory measurements. VOC species, including acetic acid, 334 propionic acid and ethanol, were also clearly elevated in the two plumes. As the aircraft 335 was further away from the CAFOs, the emissions from different sources inside CAFO 336 facilities have been well mixed prior to sampling. Thus, separation of the VOCs sources 337 from different parts of the operations occurring within the facilities is not possible using 338 aircraft measurements.

339 Figure 7 (A) shows scatterplots of acetic acid versus NH_3 from the three flights 340 over northeastern Colorado during the SONGNEX campaign. The correlation between 341 acetic acid and NH_3 is strong for all of the three flights (R=0.81-0.87). Two different enhancement ratios of acetic acid to NH₃ were observed from aircraft measurements, 342 which are close to the determined emission ratios from beef feed yards $(30.2\pm5.5\times10^{-3})$ 343 ppb/ppb) and dairy farms $(6.4\pm0.6\times10^{-3} \text{ ppb/ppb})$ from mobile laboratory measurements, 344 345 respectively. It implies that the enhancement ratios of acetic acid to NH₃ may be used as 346 an indicator for emissions from different animal types. The relative contributions to NH_3 347 enhancements between dairy and beef cattle can be estimated based on data in Figure 7A. 348 The fractional contributions to NH₃ enhancements from beef cattle are estimated in the 349 range of 0.71-0.98 based on the three SONGNEX flights (March 28: 0.98±0.01; March 29: 0.71±0.11; April 13: 0.96±0.02). Combining the three SONGNEX flights in 350 351 northeastern Colorado, beef cattle contribute 90±4% of measured NH₃ enhancements on

these flights. This evidence suggests that beef cattle are more important for NH₃ emission
from CAFOs in northeastern Colorado.

354 The enhancement ratios of other VOC species relative to NH₃ are also calculated 355 from aircraft measurements and they are compared with those from mobile laboratory 356 measurements in Figure 7B. The determined enhancement ratios of carboxylic acids and 357 alcohols compare well between aircraft and mobile laboratory measurements. The 358 enhancement ratios of acetone and acetaldehyde to NH₃ are more scattered in aircraft 359 measurements, as the agricultural plumes contributed only small enhancements of these 360 species over a high background. The enhancement ratios of phenol, cresol, CH_3SH and 361 DMS to NH₃ from aircraft measurements are lower than those from mobile laboratory 362 measurements. The measured concentrations of these species from aircraft measurements 363 were low. The signals of these species were only slightly higher than instrument noise 364 levels at higher concentrations range of NH_3 (~150 ppb, see example in Figure 6), but not 365 detectable in many plumes with lower NH₃ concentrations.

366 4. Conclusions

367 In this study, we measured downwind air to discern VOC emissions from CAFOs in northeastern Colorado using both mobile laboratory and aircraft measurements. We show 368 369 that carboxylic acids and alcohols dominate VOC emissions from CAFOs, whereas 370 sulfur-containing species and phenolic species are important to the odor activity values 371 and NO₃ reactivity of CAFO emissions, respectively. VOC compositions of CAFO 372 emissions determined from mobile laboratory and aircraft measurements are in good 373 agreement. Based on mobile laboratory measurements of CAFO sites, NH₃ emissions are 374 mainly from animals and their waste, whereas ethanol is predominately from feed storage 375 and handling. We applied a multivariate linear regression method to apportion the relative fractions from the two sources using NH₃ and ethanol as tracers. The determined 376 377 fractions between the two sources are different among various VOC species and animal 378 types. In general, phenolic species and nitrogen-containing species are mainly associated 379 with emissions from animals and their waste for the investigated CAFOs. Significant 380 contributions from feed storage and handling are observed for carboxylic acids, alcohols and carbonyls. We also proposed that ambient enhancement ratios of acetic acid to NH₃ 381 382 may be used as an indicator to separate CAFO emissions from different animal types.

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551 Figures



Figure 1. (A and B) Drive track of mobile laboratory color- and size-coded by NH_3 (A) and ethanol (B) concentrations around a beef feed yard (beef #1). The prevailing wind is shown by wind barbs (light blue flags) in the map. (C) Time series of NH_3 , CH₄, CO₂, N₂O, ethanol, acetic acid and acetone measured downwind of the beef feed yard. Numbers (1-4) in (A) and (C) are used to allow alignment of the mobile laboratory locations on the map with the corresponding time series in panel C.



Figure 2. (A and B) Drive track of mobile laboratory color- and size-coded by NH_3 (A) and ethanol (B) concentrations downwind of a dairy farm (dairy #1). The prevailing wind is shown by wind barbs (light blue flags) in the map. (C) Time series of NH_3 , CH_4 , CO_2 , N_2O , ethanol, acetic acid and acetone measured downwind of the dairy farm. Numbers (1-4) in (A) and (C) are used to allow alignment of the mobile laboratory locations on the map with the corresponding time series in panel C.

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Figure 3. The fractional contributions of different VOC classes to the total VOC
concentrations (A), odor activity values (B), OH reactivity (C) and NO₃ reactivity
(D) for the six investigated CAFO sites. The mean wind speeds during the
measurements of the CAFO sites are shown in (E). The mean values for the five

- 575 parameters from each CAFO are shown at the top of each panel.
- 576





578

579 Figure 4. The relative contributions of feed storage+handling, animal+waste and

580 milking parlors (only for dairy farm #2) to emissions of different VOC species and

- 581 total VOC for the investigated CAFO sites. DMDS: dimethyl disulfide; DMA:
- 582 dimethylamine; TMA: trimethylamine.
- 583



585 Figure 5. The relative contributions of each VOC class to emissions from feed

586 storage+handling (A), animal+waste (B) and milking parlors (C).



589 Figure 6. Time series of NH₃ and various VOC species of two agricultural plumes

590 measured from NOAA WP-3D on April 13, 2015 during the SONGNEX campaign.





594 Figure 7. (A) Scatterplot of acetic acid versus NH₃ from the three SONGNEX flights

- 595 in northeastern Colorado. The two black lines and gray-filled areas indicate
- 596 emission ratios of acetic acid to NH₃ from beef feed yards and dairy farms
- 597 determined from the mobile laboratory measurements, respectively. (B)
- 598 Comparison of enhancement ratios of VOCs to NH₃ between mobile laboratory and
- 599 aircraft measurements in northeastern Colorado.
- 600