# <sup>1</sup> Spectrometric measurements of atmospheric propane (C<sub>3</sub>H<sub>8</sub>)

2 Geoffrey C. Toon<sup>1</sup>, Jean-Francois L. Blavier<sup>1</sup>, Keeyoon Sung<sup>1</sup>, Katelyn Yu<sup>1,2</sup>

3 <sup>1</sup> Jet Propulsion Laboratory, California Institute of Technology, Pasadena, CA, 91109, USA

4 <sup>2</sup> Dept. Civil and Environmental Engineering, UC Berkeley, Berkeley, CA, 94720, USA

5 Correspondence to: Geoffrey.C.Toon@jpl.nasa.gov

6 Abstract. We report measurements of atmospheric C<sub>3</sub>H<sub>8</sub> from analysis of ground-based, solar absorption spectra

7 from the JPL MkIV interferometer. Using the strong Q-branch absorption feature at 2967 cm<sup>-1</sup>, we can measure

8 C<sub>3</sub>H<sub>8</sub> in locations where its abundance is enhanced by proximity to sources (e.g., large natural gas fields, mega-

9 cities). A case study of MkIV C<sub>3</sub>H<sub>8</sub> measurements from Ft. Sumner, New Mexico, shows, that amounts are strongly

10 correlated with ethane (C<sub>2</sub>H<sub>6</sub>), and with back-trajectories from SE New Mexico and West Texas, where the Permian

11 Basin oil and natural gas field is located. Measurements from JPL, California, also show large C<sub>3</sub>H<sub>8</sub> enhancements

12 on certain days, but more correlated with CO than C<sub>2</sub>H<sub>6</sub>. From high-altitude, balloon-borne, MkIV solar occultation

13 measurements, C<sub>2</sub>H<sub>8</sub> was not detected at any altitude (5-40 km) in any of its 25 flights,

#### 14 1. Introduction

15 Non-methane hydrocarbons such as C3H8 and C2H6 affect air quality because their oxidation enhances tropospheric

 $O_3$  and aerosol pollution. They are also sensitive indicators of fugitive losses by the oil and natural gas industry, an

- 17 important source of co-emitted methane (CH<sub>4</sub>), a greenhouse gas. These fugitive losses appear to be under-
- 18 estimated in global inventories (Dalsoren et al., 2018).

19 Atmospheric C<sub>3</sub>H<sub>8</sub> and C<sub>2</sub>H<sub>6</sub> are entirely the result of emissions at the surface. In pre-industrial times these came

- 20 from geological seeps and wild fires, but in recent times these natural sources have been surpassed by emissions
- from fossil fuel production. The latter peaked in about 1970, and then declined due to stricter regulation of

emissions from the oil and natural gas industry and automobiles. But in the past decade, this decreasing trend has reversed due to accelerated Natural Gas (NG) exploitation (Helmig et al., 2016).

24 C<sub>3</sub>H<sub>8</sub> has a lifetime of about 2 weeks in summer and 8 weeks in winter (Rosado-Reyes et al., 2007). This is mostly

- 25 dictated by how fast it is being oxidized by reactions with hydroxyl radicals and chlorine atoms. Given this 2–8
- week lifetime, a single strong source of propane has the potential to degrade air quality over most of the hemisphere.
- 27 Unprocessed, in-the-ground, "wet" natural gas is usually between 70–95% CH4, 1–15% C2H6, 1–10% C3H8, and 0–
- 3% C<sub>4</sub>H<sub>10</sub>. The latter two gases are typically extracted to form Liquified Petroleum Gas (LPG). In the northern

29 hemisphere winter, LPG contains more C<sub>3</sub>H<sub>8</sub>, while in summer it contains more butane (C<sub>4</sub>H<sub>10</sub>), reducing variations

30 in its vapor pressure.

Deleted: large variations

Deleted: amounts

Formatted: Font: Times New Roman, 10 pt

Deleted: From MkIV solar occultation measurements from balloon, C<sub>3</sub>H<sub>8</sub> was not detected at any altitude in any flight Formatted: Font: Times New Roman, 10 pt, Subscript Formatted: Font: Times New Roman, 10 pt Formatted: Font: Times New Roman, 10 pt Formatted: Font: Times New Roman, 10 pt Formatted: Font: Times New Roman, 10 pt

Moved down [1]: In contrast, the lifetime of  $C_2H_6$  is 2–6 months, which is 3–4 times longer than that of  $C_3H_8$ . Moved (insertion) [1]

**Deleted:** In contrast, the lifetime of  $C_2H_6$  is 2–6 months, which is 3–4 times longer than that of  $C_3H_8$ .

ooking, especially in Deleted: is
s increasingly
natural gas, the NG
h wet NG.
ough in situ
ppt_at Zeppelin Deleted: report
.4 ppb. These C <sub>3</sub> H <sub>8</sub> Deleted: of essentially zero
ltiple sites Helmig et
ob in winter, with <b>Deleted:</b> a
09.5–2014.5, but no
vious measurements
1) reported a 0-
al (2015) reported a
bove the
005–2010 based on
and 2014, and as a
ze that this increase
in North America,
in 1984. It covers Deleted: K
e photoconductor
•
observations a
observations a V is primarily a Deleted: K
bbservations a V is primarily a Deleted: K n flights it makes
beservations a V is primarily a n flights it makes ifferent days from 12
bbservations a V is primarily a Deleted: K n flights it makes ifferent days from 12
bbservations a     V is primarily a     Deleted: K       n flights it makes     ifferent days from 12
beservations a V is primarily a Deleted: K n flights it makes ifferent days from 12
beservations a V is primarily a Deleted: K n flights it makes ifferent days from 12

77	The analysis of the MkIV spectra was performed with the GFIT (Gas Fitting) tool, a nonlinear, least-squares,	(	D
78	spectral-fitting, algorithm developed at JPL. GFIT has been previously used for the Version 3 analysis (Irion et al.,		
79	2002) of spectra measured by the Atmospheric Trace Molecule Occultation Spectrometer, and it is currently used for	(	D
80	analysis of Total Carbon Column Observing Network (TCCON) spectra (Wunch et al., 2011) and for MkIV spectra		
81	(Toon et al., 2016; 2018a; 2018b). The entire package including spectral fitting software, spectroscopic linelists, and		
82	software to generate a priori VMR/T/P profiles, is termed GGG.		
83	GFIT scales the atmospheric gas volume mixing ratio (VMR) profiles to fit calculated spectra to those measured.		
84	For C <sub>3</sub> H <sub>8</sub> , a 5.4 cm <sup>-1</sup> -wide fitting window centered on the Q-branch at 2967 cm <sup>-1</sup> was used. The atmosphere was		
85	discretized into 70 layers of 1 km thickness. C <sub>3</sub> H <sub>8</sub> and four interfering gases (H <sub>2</sub> O, CH <sub>4</sub> , C <sub>2</sub> H <sub>6</sub> , HDO) were adjusted.		
86	Two frequency stretches were retrieved (telluric and solar). The spectral continuum was fitted as a straight line, and		
87	a zero-level offset was fitted. So that's a total of 10 simultaneously-fitted scalars. In addition, the solar pseudo-		
88	transmittance was computed (but not adjusted).		
89	The assumed temperature, pressure and H <sub>2</sub> O profiles were based on the NCEP 6-hourly analyses for solar noon of		
90	each day. The a priori vmr profiles were based on NH mid-latitude profiles. This is the same scheme as used by the	(	D
91	GGG TCCON analysis (Wunch et al., 2015), but here we apply it to the Mid-IR MkIV spectra rather than the Short-		D
92	Wave JR TCCON spectra.	(	D
93	To estimate the sensitivity of the retrieved $C_2H_{2}$ to uncertainties in the assumed a priori profiles of T/P and		
94	interfering cases (especially H $_{20}$ CH $_{10}$ ) we retrieve the post-2000 CsH $_{20}$ a second time using GGG2020 an undated		D
95	version of the GGG code with improved a priori VMR/T/P profiles based on the GGOS-EP-IT analysis (Laughner et /	1	D
96	al 2021) The results shown in figure B 2 illustrate that this changes the retrieved $C_2H_0$ by less than 10% with a	1	D
97	bias of only 1.1%.	$\leq$	
-	•		ע ק
98	2.3. Spectroscopy	$\int \lambda$	F
1		2	F
99	It is clear from the infra-req <u>tab</u> spectrum of C <sub>3</sub> H <sub>8</sub> (Fig.1), measured at <u>Pacific North-West National Laboratory</u>		
	1 = 1 = 1 = 1 = 1 = 1 = 1 = 1 = 1 = 1 =	10000	- <b>P</b>

- 100 (Sharpe et al., 2004), that the feature at 2967 cm<sup>-1</sup>, caused by various  $CH_2$  and  $CH_3$  stretch vibrational modes, is
- 101 far the strongest in the entire infrared. So for solar occultation spectrometry, this is by far the best choice For
- 102 thermal emission spectrometry from cold planets such as Titan, however, these bands are not covered by
- 103 Cassini/CIRS since the thermal Plank function of such planets weakens rapidly above 2000  $\text{cm}_{k,t}^{-1}$  Thus, the much
- 104 weaker bands below 1400 cm<sup>-1</sup> must be used (Sung et al., 2013).

# Deleted: K

Deleted: exactly	
Deleted: 2014	
Deleted: W	

Å	Deleted: instead of GGG2014						
λ	Deleted: appendix						
-(	Deleted: A						
(	Deleted: show						
-(	Deleted: rms						
Å	Deleted: lab						
λ	Formatted: Font color: Text 1						
1	Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1						
(	Formatted: Font color: Text 1						
-(	Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1						
	<b>Deleted:</b> . For thermal emission spectrometry from cold planets such as Titan, on the other hand, then the much weaker CH <sub>3</sub> deform bands around 1400 cm <sup>-1</sup> would be better (Sung et al., 2013).						
Ì	Formatted: Font: 10 pt						
Ì	Formatted: Superscript						
Ý,	Formatted: Font: 10 pt						
Formatted: Font: 10 pt, Superscript							
Ì	Formatted: Font: 10 pt						
V)	Formatted: Font: 10 pt						
Ì	Formatted: Font: 10 pt						



146	spectrometric	"cross-talk"	between thes	se two gases	should	be modest;	we con	mpute a	Pearson	Correlation	
	·										

147 Coefficient of -0.7 between the CeHs and CeHs. Further discussion on this topic can be found in Appendix A. The

upper-middle panel shows that the residuals (measured-calculated transmittance) have some systematic features of 148

~0.5% in magnitude, especially in the vicinity of the H2O line at 2966.0 cm<sup>-1</sup>. The topmost panel shows the 149

150 residuals to a fit performed without any CoHe absorption lines. It looks surprisingly similar to the fit performed with

151 C2H8 lines, such is the ingenuity of the spectral fitting algorithm in adjusting the H2O, CH4, and C2H6 to compensate

for the missing C2H8. The overall RMS residual in the no-C2H8 case is 0.3934%, as compared with 0.3658% when 152

 $C_{2}H_{8}$  is included. This is quite significant considering that residuals are dominated by the H<sub>2</sub>O line at 2966.0 cm<sup>-1</sup> 153

- 154 and are unaffected by whether CeHe is included or not. The residuals in the topmost panel (d) are larger in the
- 155 vicinity of the C<sub>3</sub>H<sub>8</sub> Q-branch, 2967-2968 cm<sup>-1</sup> than those in panel (c)

156



Figure 2. The average of 5000 ground-based MkIV spectral fits. Black diamonds represent measured spectrum. 157 158 Black line the fitted calculation. Colored lines represent the contributions of different gases. Panel (a) shows the full transmittance range. Panel (b) zooms into the 0.95–1.00 range to help see the weak absorbers (C<sub>2</sub>H<sub>6</sub>, HDO, and the 159 solar lines). Panel (c) show residuals (Measured-Calculated); these are generally below 0.5%. Panel (d) shows the 160 161 residuals when  $C_3H_8$  is excluded from the calculation.

162 Considering the weakness (and smoothness) of the C<sub>3</sub>H<sub>8</sub> Q-branch in comparison with the residuals and the 163 contributions of the other gases, we were at first skeptical that a useful C3H8 column measurement could be

Deleted	:
X	

	Deleteu:
	Formatted: Subscript
1	Formatted: Subscript
Ì	Formatted: Subscript
)	Formatted: Subscript
ì	Deleted: top
1	Formatted: Subscript
Ì	Formatted: Subscript
Ì	Formatted: Subscript
J	Formatted: Subscript
Ò	Formatted: Subscript
Ņ	Formatted: Subscript
Ņ	Formatted: Subscript
V	Formatted: Subscript
	Formatted: Superscript
	Formatted: Subscript
	Formatted: Superscript
	Formatted: Not Superscript/ Subscript
	Formatted: Not Superscript/ Subscript
1	<b>Deleted:</b> Considering the weakness (and smoothness) of the C <sub>2</sub> H <sub>2</sub> O-branch in comparison with the residuals and the

contributions of the other gases, we were at first skeptical that a useful C3H8 column measurement could be extracted from such spectral fits. But since the analysis of the MkIV spectra is highly automated, it took only a few hours to run the C3H8 window shown in Fig.2 over the 5000 MkIV ground-based spectra.

#### Deleted: → →

Deleted: Bottom p	
<b>Deleted:</b> <i>Middle p</i>	
Deleted: Top p	
Formatted: Subscript	
Formatted: Subscript	

(....[1])

182 extracted from such spectral fits. But since the analysis of the MkIV spectra is highly automated, it took only a few

- 183 hours to run the C<sub>3</sub>H<sub>8</sub> window over all 5000 MkIV ground-based spectra.
- 184

#### 185 **3. Results**

#### **186** Table 1 lists the observation sites from where MkIV has made ground-based observations up to the end of 2019. The

187 <u>vast majority are from three sites: JPL, Mt. Barcroft, and Ft. Sumner.</u>

#### 188

195

	Town	<u>State</u>	Nobs	Nday	Latitude	Longitude	<u>Altitude</u>	<b>Terrain</b>	<b>Years</b>	•	Formatted Table
					<u>(deg.)</u>	<u>(deg.)</u>	<u>(km)</u>		<b>Operated</b>		
	Esrange	Sweden	160	32	67.889	+21.085	<u>0.271</u>	Boreal	<u>1999–2007</u>		
	Fairbanks	Alaska	124	46	64.830	-147.614	0.182	Boreal	<u>1997</u>		
	Lynn Lake	<u>Manitoba</u>		_11	<u>56.858</u>	<u>-101.066</u>	<u>0.354</u>	Boreal	<u>1996</u>		Formatted: Font color: Text 1
	Mt. Barcroft	<u>California</u>	1369	<u>258</u>	37.584	-118.235	<u>3.801</u>	Alpine	<u>1994–2002</u>		
	Mtn. View	<u>California</u>	7	4	37.430	-122.080	<u>0.010</u>	<u>Urban</u>	<u>1987, 2001</u>		Formatted: Font color: Text 1
	Daggett	<u>California</u>	33		34.856	-116.790	0.626	Desert	<u>1993</u>		
	Ft. Sumner	New Mex.	521	<u>106</u>	34.480	-104.220	1.260	Steppe	<u>1989–2019</u>		
	TMF	California	475	_45	34.382	-117.678	<u>2.257</u>	Alpine	<u>1986–2009</u>		
	JPL (B183)	California	2273	<u>690</u>	34.199	-118.174	0.345	Urban	1985-2020		
	JPL (mesa)	<u>California</u>		5	34.205	<u>-118.171</u>	<u>0.460</u>	<u>Urban</u>	<u>1988–1989</u>		Formatted: Font color: Text 1
	Palestine	Texas	4	3	<u>31.780</u>	<u>-95.700</u>	<u>0.100</u>	Rural	<u>1989</u>		Formatted: Font color: Text 1
	<u>McMurdo</u>	Antarctica	37		<u>-77.847</u>	+166.728	<u>0.100</u>	Polar	<u>1986</u>		Tormatted. Font color. Text T
189											
190	Table 1., The tv	<u>velve sites fro</u>	<u>m where</u>	<u>MkIV h</u>	<u>as made gro</u>	und-based obs	ervations, al	ong with the	<u>e number of</u>		Formatted: Font: Bold, Italic
191	observations an	nd observation	<u>n days fro</u>	om each	site, years o	f operations, <u>t</u> l	heir location	and terrai	1 type. They gr	<u>eved</u>	Formatted: Font: Italic
192	out sites have th	he fewest obse	ervations	(only 1	% of total) a	nd are not incl	uded in the H	igures 5-7	and A.1 to red	uced	Formatted: Font: Italic
193	color ambiguity	2.									Formatted: Font: Italic
											Formatted: Font: Italic
194	Fig. 3 shows M	kIV ground-b	based C <sub>3</sub> F	I8 colun	nns, color co	ded by site alti	tude. The da	ta were filte	red: only poin	ts	

with uncertainties $< 1.5 \times 10^{16}$ were plotted, reducing the number of plotted points from 5000 to 4700. The top panel	
(a) shows that at the high-altitude sites (Mt. Barcroft at 3.8 km is Red: Table Mountain Facility at 2.26 km is	

196	(a) shows that at the high-altitude sites (Mt.	t. Barcroft at 3.8 km is Red; Table Mountain Facility at 2.2 <u>6 km is</u>
-----	--	---

197 Orange) the retrieved  $C_3H_8$  columns are centered around zero. Also, the data acquired in Sep 1986 from 0.1 km in

198 Antarctica (dark blue) are centered around zero. Data acquired from Ft. Sumner, NM, at 1.2 km (lime) have large

variations, from zero to nearly  $8 \times 10^{16}$  molecules.cm<sup>-2</sup>, as do the data from JPL at 0.35 km (cyan). Other sites with detectable C<sub>3</sub>H<sub>8</sub> include Daggett, CA, (0.6km), Esrange, Sweden (0.26km) in the winter, Fairbanks, AK (0.2km),

Deleted: 5E+	
Deleted: =	
Deleted: =	
Deleted: also	
Deleted: E+	

Moved (insertion) [2]
<b>Moved up [2]:</b> So C <sub>3</sub> H <sub>8</sub> has only been measured by MkIV from northern hemisphere sites within the PBL.
Formatted: Superscript
Deleted: , probably
Deleted: ing

and Mountain View, CA in late 1991. So C<sub>3</sub>H<sub>8</sub> has only been measured by MkIV from northern hemisphere sites

<sup>202</sup> within the PBL. Panels (b) and (c) show the same  $C_3H_8$  columns, but plotted versus year and day.

High C<sub>3</sub>H<sub>8</sub> values (>4x10<sup>16</sup> molecules.cm<sup>2</sup>) can occur at any time of year at JPL (cyan) but most commonly in late

summer, as is the case for other pollutants, e.g. CO. This reflects the meteorology (stagnant conditions in the LA

- basin in summer with little replacement of polluted air with clean air from outside). Averaging kernels for these
- 215 C<sub>3</sub>H<sub>8</sub> measurements are discussed and illustrated in Appendix B. Suffice it to say here that they range from 0.9 to
- 216 1.4 and increase with altitude.





218	Figure 3. MkIV C <sub>3</sub> H <sub>8</sub> column abundances from <u>8 out of</u> 12 sites, color-coded by site altitude, as illustrated in panel
219	(a): Violet=0.1_km (McMurdo); dark blue=0.18 km (Fairbanks); light blue=0.27km (Esrange); cyan=0.35 km
220	(JPL); Green=0.63 km (Daggett); lime=1.2 km (Ft. Sumner); orange=2.26 km (TMF); red=3.8 km (Mt. Barcroft).
221	The reported uncertainties in our C3H8 column measurements are based on the rms fitting residuals compared with
222	the sensitivity of the spectrum to $C_3H_8$ (Jacobians). At the highest site, Barcroft at 3.8 km (P=0.65 atm.), where the
223	interfering H2O and CH4 absorptions are relatively weak and narrow, the C3H8 column uncertainties are generally
224	smaller than 1015 molecules.cm <sup>-2</sup> . But since the columns themselves are even smaller, no C <sub>3</sub> H <sub>8</sub> is detected at
225	Barcroft. At the lower altitude sites such as JPL and Ft. Sumner, the increased interference from H <sub>2</sub> O and CH <sub>4</sub> cause
226	the C3H8 column uncertainties to be much larger, generally around 5x1015 molecules.cm-2 at low airmass and
227	worsening rapidly toward higher airmasses. But the C2H8 increases far more, allowing, C3H8 to be detected at these
228	low-altitude sites under polluted conditions, despite the poorer absolute uncertainties.

Deleted: all Deleted: blue Deleted: 0 Deleted: 5

(	Deleted: This
·····(	Deleted: s
$\supset$	Formatted: Subscript
) (	Formatted: Subscript

236 High C<sub>3</sub>H<sub>8</sub> values are also seen at Ft. Sumner, NM (lime), especially in recent years. This was initially a surprise to

237 us because this area has a very low population density, so we naively assumed that we would be measuring

238 background levels of atmospheric pollutants here.

239 We know that the apparent variations in  $C_3H_8$  are real, rather than artifacts, from their strong correlation with  $C_2H_6$ .

Figure 4 compares column-averaged  $C_3H_8$  mole fractions (top panels) with those of  $C_2H_6$  (bottom panels), the latter

241 retrieved using different spectral lines than those shown in Fig.2. These are the same total C<sub>3</sub>H<sub>8</sub> columns shown in

Fig.3, but divided by the total column of all gases, which is inferred from the surface pressure. The resulting

243 column-average mole fractions, denoted Xgas, are less sensitive to the site altitudes being different and more easily

244 compared with in situ measurements being in units of mole fraction.

245 The upper and lower rows of Fig.4 shows the XC3H8 and XC2H6 time series, respectively, plotted versus year (left)

and versus day of the year (right) The data were filtered such that only points with  $XC_3H_8$  uncertainties  $\leq 0.74$  ppb

and  $C_2H_6$  uncertainties < 0.10 ppb were plotted. This reduced the total number of points from 5000 to 4700, so only

- 248 the best 94% of the data are plotted. It is clear that at JPL (cyan) C<sub>3</sub>H<sub>8</sub> has decreased since the 1990s, but that at Ft.
- 249 Sumner (lime) it has increased over the past decade. The data from these two sites will be explored later.





Figure 4. Top panels show measurements of the column-averaged C<sub>3</sub>H<sub>8</sub> mole fractions (XC<sub>3</sub>H<sub>8</sub>). Bottom panels
 show XC<sub>2</sub>H<sub>6</sub>. Left panels show the variation with year. Right-hand panels show the seasonal variation. <u>Points are</u>
 <u>color-coded by observation site altitude\_as in Fig.3.</u>





Formatted: Font: Times New Roman, 10 pt

Formatted: Font: Times New Roman, 10 pt





263  $\int_{C_2H_4}$  is four times longer-lived than C<sub>3</sub>H<sub>8</sub> and never goes to zero because there is always a substantial free

tropospheric C<sub>2</sub>H<sub>6</sub> component, even in the SH, which varies seasonally: high in spring, low in fall. The Antarctic

265 measurements (blue) are very low (0.2-0.3 ppb) and most probably even lower during the rest of the year, because

days 250 to 300 represent the springtime peak, not the fall. The highest C<sub>2</sub>H<sub>6</sub> ever, measured from JPL (cyan) was in

late 2015 (day 314) as a result of the Aliso Canyon natural gas leak (Conley et al., 2016). This event is further

268 discussed later and also in Appendix C.

269 Figure 5 shows the XC<sub>2</sub>H<sub>6</sub>/C<sub>3</sub>H<sub>8</sub> correlation plot for all sites. This uses the exact same data, filtering, and color-

 $\label{eq:scheme as for Fig. 4. At JPL (cyan) the correlation is positive but weak. At Ft. Summer, there are episodes of both$ 

271 gases being enhanced with a strong correlation. In fact, the highest VMRs of C<sub>2</sub>H<sub>6</sub> were seen from Ft. Sumner, even

more than from JPL during the Aliso Canyon gas leak in late 2015.



Deleted:	The lower panels of Fig.4 show XC <sub>2</sub> H <sub>6</sub> . Th
Deleted:	is
Formatt	ed: Subscript
Formatt	ed: Subscript
Deleted:	that
Formatt	ed: Font: Times New Roman, 10 pt
Deleted: than they	The Antarctic measurement (blue) are even lower appear because
Deleted:	in Antarctica
Deleted:	ever
Deleted:	was
Deleted:	there



274 *Figure 5.* The correlation between XC<sub>2</sub>H<sub>6</sub> and XC<sub>3</sub>H<sub>8</sub> for all sites, color-coded by site altitude <u>as in Fig.3</u>.

#### 275 <u>3.1. Averaging Kernels.</u>

273

- 276 Figure 6 shows all kernels for the 5000 measurements presented in this paper, color-coded by site altitude (red=3.8
- km; orange=2.2 km; lime=1.2 km; cyan=0.35 km; blue < 0.2 km) as in the main body of the paper. The kernels
- increase with altitude but with <40% variation over the 0-30 km altitude range. Note that the kernels representing
- the 3.8 km site begin at P=0.7 atm. And the kernels representing the 2.2 km site begin at P=0.8 atm.



298 these are smaller effects than those of site altitude or SZA.



**300** *Figure 7. A priori* C<sub>3</sub>H<sub>8</sub> *profile used in these retrievals.* 



320	Figure & XC3H8, XC2H6 and XCO at Ft. Sumner. Since all the observations are made from the same altitude, it no		Moved down [5]: MKIV didn't visit Ft. Sumner from 1997
321	longer makes sense to color code by site altitude. So instead we color-code by mean bearing of the back-trajectory		to 2004 because it was performing high-latitude balloon flights from Alaska and Sweden.
322	over the previous 36 hours. Dark blue=30°; Light blue =90°, Cyan=120°; Green=180°; Lime=220°; Orange=	Y	Deleted: 6
323	300 ° Red=350 °. MKIV didn't visit Ft. Sumner from 1997 to 2004 because it was performing high-latitude balloon		Moved (insertion) [5]
324	flights from Alaska and Sweden.	(	Deleted: ,
325			
326	Figure & shows that between 1990 and 2005 there was a decrease in C2H6 and C3H8 measured in Ft. Sumner, by		Deleted: 6
327	about a factor 2 over 15 years. In recent years (since 2014), however, there has been a large increase in C2H6 and		
328	C <sub>3</sub> H <sub>8</sub> measured at Ft. Sumner, but only when the wind direction is from the SE quadrant (green-cyan, colors). We see		Deleted: /lime
329	no increase associated with other wind directions (red, blue, orange, yellow, lime).		
330	At Ft. Sumner CO has no correlation with wind direction, nor with C2H6 or C3H8. The majority of days have a		
331	column average CO of 75±10 ppb. But there are occasional enhancements up to 120 ppb, likely due to large but		
332	distant fires. We do not pursue the Ft. Sumner CO data any further, beyond proving that the C3H8 sources are		Deleted: . They are of no value
333	different from those of CO.	(	Deleted: other than
334	CH <sub>4</sub> is also measured by MkIV. Over the 30-year measurement period XCH <sub>4</sub> has grown from 1650 to 1850 ppb.		
335	This secular increase is much larger than any variation due to wind direction. So to be useful, the CH4 data would		
336	have to be detrended, which is not simple given its non-linear growth. Even within the past 4 years, the correlation		
337	of XCH4 with XC3H8 was very weak. This is to be expected since the background abundance of CH4 is more than		
338	1000x larger than C <sub>3</sub> H <sub>8</sub> , whereas wet NG is only 6 times richer in CH <sub>4</sub> than C <sub>3</sub> H <sub>8</sub> (in the Permian basin). So the NG-	(	Deleted: from the Permian Basin
339	induced enhancement of CH4, as a fraction of its atmospheric background level, will be much, smaller than that of	(	Deleted: .
340	$C_3H_8$ .	(	Deleted: be more than 100 times
1			
341	Figure <u>9</u> shows a XC <sub>3</sub> H <sub>8</sub> -XC <sub>2</sub> H <sub>6</sub> scatter plot using just the Ft. Sumner data. Error bars are much larger for XC <sub>3</sub> H <sub>8</sub>		Deleted: 7
342	than for XC <sub>2</sub> H <sub>6</sub> . This is because the C <sub>2</sub> H <sub>6</sub> transitions are stronger and form narrower features, both of which make		
343	the retrievals more precise and definitive, whereas most of the C3H8 absorption is smeared into a broad continuum		
344	which provides little information for a retrieval in which the continuum level is fitted. The C2H6 features used in the		Deleted: this type of
345	actual $C_2H_6$ retrieval are at 2976.6 and 2986.6 cm <sup>-1</sup> (not shown) and are 3–4 times stronger than those seen in Fig.2.		







384 Figure <u>10</u>, Hourly locations for the back-trajectories, color-coded by <u>retrieved XC3H8</u>, Blue=0 ppb, Green=2 ppb; Red=4 ppb. Trajectories for which the  $XC_3H_8$  uncertainty exceeded 0.74 ppb are excluded, resulting in only 373 out 385 of 520 trajectories being shown. Ft. Summer lies at 34.2N, 104.2W, close to the center of the figure at the confluence 386 387 of all the back-trajectories. Each point represents a 1-hour time step, so that the wind speed is apparent from the 388 separation of points. Winds from the West are typically stronger than those from the SE quadrant. Trajectories are 389 underlaid by a map of New Mexico and neighboring states. The Permian Basin, encircled by the thick purple line, 390 underlies SE New Mexico and much of West Texas. Many of the trajectories from the SE have spent 30+ hours over 391 the Permian Basin.

We also made a scatter plot for CO (not shown) but there was no correlation between CO and wind direction, or between CO and  $C_3H_8$ . This rules out the possibility that the enhanced  $C_3H_8$  and  $C_2H_6$  were somehow associated with distant urban pollution or wild fires.

 $\label{eq:295} This result leads to speculation on what might be enhancing C_2H_6 and C_3H_8 when the winds come from the SE$ 

sector. One of the biggest natural gas production fields in the US lies in the Permian Basin, which underlies the

397 South-East corner of New Mexico and West Texas, as illustrated in Figure <u>10</u>, This region also includes processing

398 plants where the heavier gases are stripped out of the wet NG, storage facilities for the resulting Natural Gas Liquids

399 (LPG+ethane+pentane), and pipelines. <u>The Permian Basin is by far the largest "liquids-rich" (rich in heavy</u>

400 hydrocarbons) gas field in the USA (https://www.spglobal.com/platts/plattscontent/\_assets/\_images/latest-

Deleted: 8 Deleted: . Deleted: . Deleted: nd

408 news/20191219-rig-count.jpg). This would suggest that the enhanced C<sub>2</sub>H<sub>6</sub> and C<sub>3</sub>H<sub>8</sub> is the result of losses from NG

409 production, although this cannot be proven with just one instrument at one site. We would need instruments upwind

410 and downwind to make an accurate assessment of the fluxes.



#### 411 412

412		ananana a
413	Figure 11; NG production in the lower 48 states of the USA in 2009. Data from the Energy Information	1
414	Administration: <u>https://www.eia.gov/oil_gas/rpd/conventional_gas.pdf.</u> Superimposed are the locations (purple	
415	pentangular star) of the four sites discussed in detail in this paper: Ft. Sumner in Eastern NM is labelled "FTS". The	1
416	JPL site in California is labelled "JPL". The locations of the NOAA sites in Utah (UTA) and Oklahoma (SGP) are	1
417	also included. The Permian basin lies in the SE corner of NM and West Texas <sub>7</sub>	Ι,
418	The Permian basin currently produces 16 billion cu.ft./day of NG	1
419	(https://www.eia.gov/petroleum/drilling/pdf/permian.pdf) over an area of 220,000 km <sup>2</sup> . The molar volume of an	
420	ideal gas at STP is 22.4 liters. One cu. ft. is 28.3 liters. So 16 billion cu. ft. is 20 billion moles of NG or 120x10 <sup>32</sup>	Witness.
421	molecules per day. Over an area of 220,000 km <sup>2</sup> or 2.2x10 <sup>15</sup> cm <sup>2</sup> , this represents an average areal production of	Statements.
422	55x10 <sup>17</sup> molec./cm <sup>2</sup> /day. Assuming that the Permian basin is 480 km wide, at an average low-level wind speed of 15	1
423	km/hour, an air parcel will take 32 hours (1.33 days) to traverse the Basin, during which time 73x10 <sup>17</sup>	
424	molecules/cm <sup>2</sup> will have been extracted. Of this, 10% will be C <sub>3</sub> H <sub>8</sub> (Howard et al., 2015), so if all this production	
425	were released into the atmosphere we would expect a C <sub>3</sub> H <sub>8</sub> column enhancement of 73x10 <sup>16</sup> .	
426	In airmasses with trajectories from the SE, we see maximum $C_3H_8$ column enhancements of only $3x10^{16}$	
427	molecules/cm <sup>2</sup> , which suggests that only 4% of the NG escapes into the atmosphere and that 96% of the NG is	ll
428	successfully captured (or burnt by flaring).	
		b
429	In the Permian Basin, NG is 13.7% C <sub>2</sub> H <sub>6</sub> and yet the observed ethane enhancements are slightly smaller than those	ų,
430	of C <sub>3</sub> H <sub>8</sub> suggesting that only ~3% of the NG escapes. Assuming a 3% leak rate, there will also be an enhancement	K
431	of CH4 of about 14x10 <sup>16</sup> molec.cm <sup>-2</sup> , but this represents only 0.4% of the total CH4 column above Ft. Sumner and	1
432	will therefore be difficult to discern in the presence of other confounding factors (stratospheric transport, varying	and the second second

433 tropopause altitude, seasonal and longer-term changes). Of course, all this analysis assumes that the Permian basin

## Deleted: 9

# Deleted: (a)

#### Deleted:

(b): [Temporarily removed -- awaiting permission] Illustrating the high number of "liquids-rich" drilling rigs in the Permian Basin, as of Dec. 2019, underscoring itsdominance for propane production in the USA. From https://www.spglobal.com/platts/en/market-insights/latestnews/natural-gas/121919-us-oil-gas-rig-count-rises-forsecond-straight-week-enverus¶

Formatted: Font: Times New Roman, 10 pt

Formatted: Line spacing: 1.5 lines

Formatted: Font: Times New Roman, 10 pt

Formatted: Font: Times New Roman, 10 pt

#### Formatted: Font: 10 pt

**Deleted:** In recent years, the Permian basin has been producing ~15 billion cu.ft. of natural gas (NG) per day (https://www.eia.gov/petroleum/drilling/pdf/permian.pdf). A back-of the-envelope estimate of the contribution of this to the observed  $C_3H_8$  is now performed. We assume that this NG production is distributed over an area that is 160 km across. At a wind speed of 20 km/hour, an airmass will take 8 hours to traverse the gas field, during which time 0.72E+19 molecules.cm<sup>2</sup> of NG will have been extracted. Howard et al., (2015), measured the composition of NG from the Permian basin and found that it is very rich in heavy hydrocarbons, being 66.6% CH4, 13.7% C<sub>2</sub>H<sub>6</sub> and 10.3% C<sub>3</sub>H<sub>8</sub> by volume. If 4% of this were lost to the atmosphere, and 10.3% of this is C<sub>3</sub>H<sub>8</sub>, the total propane column will be enhanced by 3E+16 molecules.cm<sup>-2</sup>, which is close to that seen in the highest cases. For

#### Deleted: C2H6,

9	Formatted: Subscript
2	Formatted: Subscript
	Deleted: an
	Deleted: of 4E+16
	Deleted:
~	<b>Deleted:</b> would be expected for such a back-trajectory, which is somewhat higher than measured
	Deleted: T
- N.	

Deleted: 9E+

#### 469 is a uniform emitter and that the back trajectory wind speeds are accurate. There are likely hot spots with higher-

470 <u>than-average emissions, and regions with little NG production.</u>

# 471 A puzzle in our findings is that when both $C_3H_8$ and $C_2H_6$ are elevated, we measure 22% more $C_3H_8$ than $C_2H_6$ (see

472 fig.9). Yet independent essays of well-head wet NG find <u>33%</u> more C<sub>2</sub>H<sub>6</sub> than C<sub>3</sub>H<sub>8</sub> in the Permian basin (Howard et

473 al., 2015). So we have a 55% discrepancy. We note that the C<sub>2</sub>H<sub>6</sub> averaging kernel is 0.7 at the surface versus 0.9 for

474 C<sub>3</sub>H<sub>8</sub> (see Appendix B). So when these gases exceed their priors in the PBL, which is likely at high enhancements,

both will be under-estimated, but  $C_2H_6$  more so than  $C_3H_8$ . So this effect would cause the  $C_2H_4/C_2H_6$  ratio to be 28%

476 high, which explains half the 55% problem, Another possibility is that the C3H8 coming from fugitive, wet NG is

**477** augmented by leaks of LPG, stripped from wet NG. This would further enhance the  $C_2H_8$  (and  $C_4H_{10}$ ) with little

478  $\underline{C_2H_6}$  increase. Alternatively, there could be a systematic over-estimate of the MkIV  $C_3H_8$  due to a mundane

479 multiplicative bias in the  $C_3H_8$  spectroscopy. This would over-estimate all the  $C_3H_8$  measurements without

480 degrading the strong correlation with C<sub>2</sub>H<sub>6</sub>, but seems unlikely.

#### 481 3.3. Case Study: Ground-based measurements from JPL

482 The Jet Propulsion Laboratory (34.2N; 118.17W; 0.35 km altitude) lies at the Northern edge of the Los Angeles

483 basin. When winds are from the North (rare in summer) air quality is good. When conditions are stagnant (common

484 in summer) pollutants accumulate and so air quality is poor. C<sub>3</sub>H<sub>8</sub> measured at JPL exhibits very different behavior

to that at Ft. Sumner. It decreases over time, exhibits little correlation with C<sub>2</sub>H<sub>6</sub>, and positive correlation with CO.

486 Figure 12, illustrates these behaviors.

487 The left-hand panels of Fig. 12, shows XC3H8 time series measured from JPL, color coded by CO. The upper-left

488 panel shows a large decrease in C<sub>3</sub>H<sub>8</sub> from 1–3 ppb in 1990 to less than 1 ppb in 2019. This mirrors the decrease in

489 CO over JPL (not shown) over the same period. The lower-left panel shows a large seasonal component to the

 $C_3H_8$ , with a peak in late summer, when the air is most stagnant over JPL allowing pollutants to accumulate. The

491 highest C<sub>3</sub>H<sub>8</sub> values appear red or orange (high CO), while the lowest appear blue (low CO), implying an

association with CO. This is confirmed in the upper-right panel which plots  $C_3H_8$  directly against CO. The right-

 $\label{eq:493} \mbox{hand panels are color-coded by year. The $C_3H_8$ correlation is mostly a result of both gases having decreased over the $C_3H_8$ correlation is mostly a result of both gases having decreased over the $C_3H_8$ correlation is mostly a result of both gases having decreased over the $C_3H_8$ correlation is mostly a result of both gases having decreased over the $C_3H_8$ correlation is $C_3H_$ 

494 30-year record. But even within each year, there still remains a positive correlation. This does not necessarily mean

495 that C<sub>3</sub>H<sub>8</sub> and CO have the same source, but that their sources are spatially coincident.

# **Deleted:** Of course, in cases of higher wind-speeds, or trajectories that partially circumvent the basin, the duration will be less than 8 hours and so the uptake of hydrocarbons will be smaller.

#### Deleted: .

Deleted: always

**Moved down [4]:** One possibility is that the  $C_3H_8$  coming from leaking wet NG is augmented by leaks of LPG, stripped from wet NG. This would further enhance the  $C_3H_8$  (and  $C_4H_{10}$ ) with little  $C_2H_6$  increase.

#### Deleted: ¶

Formatted: Not Superscript/ Subscript

Deleted: /C<sub>3</sub>H<sub>8</sub> Deleted: 5

Deleted: low Deleted: Deleted: m, but not all Moved (insertion) [4] Deleted: One

Deleted: leaking

Deleted: ¶

# Deleted: 2

Deleted: 0 Deleted: 0



519 520 521

*Figure 12,* Column-average  $C_3H_8$  above JPL. Left Panels: The time series color-coded by CO (red=250 ppb; green=130 ppb; blue=100 ppb). Right Panels: The relationship between  $XC_3H_8$  and CO and  $C_2H_6$  color-coded by year (blue=1990; green=2005; red=2019).

Deleted: 0

 $\label{eq:constraint} 524 \qquad \text{The lower-right panel shows $C_3H_8$ plotted versus $C_2H_6$. There is a weak correlation at JPL. The high $XC_2H_6$ values $V_1 = V_1 = V_2 =$ 

525 exceeding 2.0 ppb were measured in day 314 of 2015 when JPL was downwind of the Aliso Canyon NG leak.

526 Appendix C shows a HYSPLIT back-trajectory confirming this assertion. This spike can also be seen in Fig. 3.

527 There is no C<sub>3</sub>H<sub>8</sub> enhancement associated with the C<sub>2</sub>H<sub>6</sub> spike, since processed NG was leaking from an

528 underground storage facility, the heavy hydrocarbons (e.g., C<sub>3</sub>H<sub>8</sub>, C<sub>4</sub>H<sub>10</sub>) having already been stripped out. A 2%

529 increase in column-averaged CH4 was also noted in the plume of the Aliso Canyon leak, as shown in Appendix C.

530 California accounts for less than 1% of total U.S. natural gas production and this has declined over the past three

 $\label{eq:constraint} {\tt 531} \qquad {\tt decades (https://www.eia.gov/state/analysis.php?sid=CA).} \ {\tt Although there is natural gas extraction in the LA basin,}$ 

 $\label{eq:sigma} \text{532} \qquad \text{this is a small source compared with the Permian basin. The local natural gas is only 3% C_2H_6 and 0.3\% C_3H_8,$ 

533 (https://www.socalgas.com/stay-safe/pipeline-and-storage-safety/playa-del-rey-storage-operations) and so cannot

account for the approximately equal amounts of these gases measured at JPL by the MkIV. We speculate that the

535 C<sub>3</sub>H<sub>8</sub> measured at JPL comes mainly from LPG (e.g., used in "clean" commercial vehicles, BBQ grills, external

 $\label{eq:sigma} \text{batters, etc.}). \ \text{We can certainly rule out the possibility that the $C_3H_8$ measured at JPL is the result of wild fires, since $P_1 = 1$ and $P_2 = 1$ and $P_$ 

537 these have increased in recent years whereas the  $C_3H_8$  has decreased.

539	3.4, Comparison with In Situ Measurements	(	Deleted: 3
540	First it should be pointed out that the column-average mole fractions that are derived from the column measurements		Deleted: First a caveat:
541	will under-estimate the gas amount in the PBL for gases like C2H6 and C3H8 that reside mainly in the PBL. For		
542	example, if C3H8 resides entirely between 1000 and 800 mbar, with none in the free troposphere or stratosphere,		
543	then the column-average values will be 5 times smaller than the actual mole fractions in the PBL. So direct		
544	comparisons of the remote and in situ mole fractions should be avoided. But their behavior as a function of year or		
545	season, or gas-to-gas correlations, can still be meaningfully compared. This effect is in addition to the effect of their		
546	averaging kernels being less than 1.0 at the surface, which was discussed earlier.		
547	In situ C3H8 and C2H6 mole fractions from the Wendover, Utah (UTA) and Southern Great Plains, Oklahoma (SGP)		
548	sites were downloaded from the NOAA Global Monitoring Laboratory website:		
549	(https://www.esrl.noaa.gov/gmd/dv/data/). These sites are the closest to Ft. Sumner. These are surface flask		
550	measurements covering the period 2006 to 2017. Figure 13 illustrates these data as a function of the year (left		Deleted: 1
551	panels), the day of the year (middle panels), and the $C_3H_8-C_2H_6$ relationship (right panels). The upper panels cover		
552	the UTA site and the lower panels the SGP site. Note the factor 10 change in the y-scale: there is 10x more of these		
553	gases at SGP than at UTA. Looking at the map in Fig. 11, this is clearly because SGP lies immediately downwind of		Deleted: 9
554	the Anadarko Basin oil and NG fields under the prevailing WSW winds. In contrast, the UTA site has no major up-		
555	wind source.		

- $\label{eq:solution} 556 \qquad These in situ measurements confirm that C_3H_8 is highly variable with large enhancements being associated with oil$
- and NG production fields. At SGP the  $C_3H_8/C_2H_6$  ratio is about 0.65. This is smaller than those measured by the
- 558 MkIV, but NG in the Permian basin is much wetter (richer in  $C_3H_8$ ) than in the Anadarko basin.



**Figure 13** In situ flask measurements of  $C_3H_8$  (red) and  $C_2H_6$  (blue) from the NOAA ESRL GMD dataset (Helmig et al., 2017). Top panels show results from the UTA site and lower panels from SGP. Note the factor 10 change in the y-scale between the two sites. Left panels plot data versus year to illustrate secular trends. Middle panels versus Day of year to more clearly see the seasonal cycle. Right panels plot  $C_3H_8$  versus  $C_2H_6$ .

#### 9 3.5. Balloon Results

We also attempted to retrieve C<sub>3</sub>H<sub>8</sub> from MkIV balloon solar occultation spectra. It was not detected in any flight, despite a very good sensitivity of 0.05 ppb above 5 km. This confirms that the C<sub>3</sub>H<sub>8</sub> detected in ground-based measurements, reaching column average mole fractions of up to 4 ppb, resides mostly in the PBL. The balloon launches are typically performed only under stable, quiescent, meteorological conditions with light surface winds. Such conditions preclude uplift of air from the PBL into the free troposphere, so that C<sub>3</sub>H<sub>8</sub> stays confined to the PBL, which is opaque in limb paths due to aerosol, and so cannot be probed in occultation. This does not preclude C<sub>3</sub>H<sub>8</sub> getting up into the free troposphere at other times or in other places.

577

578

Deleted: 4

#### 581 4. Summary and Conclusions

582 We report measurements of atmospheric  $C_3H_8$  by solar absorption spectrometry in the strong Q-branch region at

583 2957 cm<sup>-1</sup>, using high resolution IR spectra from the JPL MkIV interferometer. To the best of our knowledge, these

are the first remote sensing measurements of atmospheric  $C_3H_8$ . The minimum detectable abundance is about  $10^{16}$ 

 $\label{eq:constraint} \text{molecules.cm}^{\text{-2}}\text{, which is roughly equivalent to a column average mole fraction of 0.5 ppb. This allows C_3H_8 to be$ 

measured in locations where its abundance is enhanced by proximity to sources (e.g., large gas fields, mega-cities),

587 but not in clean locations (e.g. above the PBL or away from sources). We encourage such NDACC and TCCON

stes to examine their datasets for Calla. Future improvements to the spectroscopy of the interfering gases, e.g. H2O,

589 CH<sub>4</sub>, C<sub>2</sub>H<sub>6</sub>, and other CH-containing gases currently missing might even provide for the detection of C<sub>3</sub>H<sub>8</sub> from

590 clean sites at background levels, allowing it to become a routine product of the NDACC and TCCON networks.

A case study of ground-based MkIV measurements from Ft. Sumner, New Mexico, shows, increasing C<sub>3</sub>H<sub>8</sub> and C<sub>2</sub>H<sub>6</sub>

amounts in the past decade on days when back-trajectories <u>came</u> from SE New Mexico and West Texas, where the

 $\label{eq:permian} \begin{array}{l} \mbox{Permian Basin oil and gas field is located. A case study of $C_3H_8$ measured at JPL shows a long-term decrease since} \end{array}$ 

1990 by more than a factor 2. It also shows a strong correlation with CO, a tracer of urban pollution. There is no

significant correlation between  $C_3H_8$  and  $C_2H_6$  at JPL.

596 The MKIV measurements in the case studies are not particularly useful for determining the long-term global trends

597 in C<sub>3</sub>H<sub>8</sub> or C<sub>2</sub>H<sub>6</sub>, due to their close proximity to strong sources. In the case of the Ft. Sumner the source is the

598 Permian Basin. In the case of JPL the source is the Los Angeles urban area with a population of ~15M. These

sources cause large meteorology-driven fluctuations that mask the longer-term trends.

From balloon measurements in solar occultation, propane was analyzed using the same window as for the ground-

based measurements. It was not detected at any altitude in any of our 25 flights, despite a 0.05 ppb detection limit.

This is presumably because under the stable atmospheric conditions that allow balloon launches, C<sub>3</sub>H<sub>8</sub> stays

603 confined to the PBL, which is opaque in the limb viewing geometry and so cannot be probed.

604 Appendix A: Correlations between retrieved parameters

We compute Pearson Correlation Coefficients (PCC) from the a posteriori covariance matrix for each of the 5000

506 spectral fits. The figures on the upper-left show the PCC between retrieved C<sub>3</sub>H<sub>8</sub> and C<sub>2</sub>H<sub>6</sub>. Points are plotted versus

 $\frac{1}{2}$  year with the same site-altitude-dependent coloring as in the other figures. The PCC between C<sub>3</sub>H<sub>8</sub> and C<sub>2</sub>H<sub>6</sub>

averages about -0.7, which means that they are fairly strongly anti-correlated. This is due to their overlapping

absorption features at 2967.5 cm<sup>-1</sup>. So as retrieved  $C_2H_6$  increases, retrieved  $C_3H_8$  will decrease, and vice versa. The

610 PCCs are closer to zero for the high-altitude sites (red & orange), presumably due to the reduced pressure

611 broadening and H<sub>2</sub>O causing the C<sub>2</sub>H<sub>2</sub> and C<sub>2</sub>H<sub>6</sub> absorption features to become more distinct. This anti-correlation

- **612** could be reduced by use of a wider window to introduce additional  $C_2H_6$  features that don't correlate with  $C_2H_6$  but
- 613 this would also encompass large residuals without adding any C<sub>3</sub>H<sub>8</sub> information.

Formatted: Subscript Formatted: Subscript

Deleted: a

Formatted: Font: Times New Roman, Not Italic, Font color: Text 1 Formatted: Line spacing: 1.5 lines Formatted: Font: Times New Roman, Not Italic, Font color: Text 1

Formatted: Subscript

Formatted: Font: Times New Roman, Not Italic, Font color: Text 1

Formatted: Font: Times New Roman, Not Italic, Font color: Text 1

Deleted: overlap the

Deleted: Q-branch



Formatted: Font: Bold, Font color: Text 1
Formatted: Line spacing: 1.5 lines
Formatted: Font color: Text 1
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1
Formatted: Font color: Text 1
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1
Formatted: Font color: Text 1, Subscript
Formatted: Font color: Text 1
Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1
Formatted: Space After: 6 pt, Line spacing: 1.5 lines
Formatted: Font: 10 pt, Not Italic, Font color: Text 1
Formatted: Font: 10 pt, Not Italic, Font color: Text 1, Subscript
Formatted: Font: 10 pt, Not Italic, Font color: Text 1
Formatted: Font: 10 pt, Not Italic, Font color: Text 1, Subscript
Formatted: Font: 10 pt, Not Italic, Font color: Text 1
Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1
Formatted: Font: 10 pt, Not Italic, Font color: Text 1
Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1
Formatted: Font: 10 pt, Not Italic, Font color: Text 1
Formatted: Font: Times New Roman, 10 pt, Not Italic, Font color: Text 1
Formatted: Font color: Text 1
Deleted: A
Deleted: exactly
Deleted: 2014
Deleted: 06











#### 672 Appendix C: - Aliso Canyon Underground Storage Facility: Gas Leak in late 2015

673 Aliso Canyon Underground Storage Facility is located 30 km NW of JPL. According to the Jan 4, 2016, 674 Los Angeles Times, NG leak began Oct 23, 2015 and peaked on Nov 28 at 60 Tons of CH4 per hour. By Dec 22 leak rate had decreased to 30 Tons per hour as the underground storage pressure dropped from the initial 2700 psi. 675



Deleted: Appendix B: C<sub>3</sub>H<sub>8</sub> and C<sub>2</sub>H<sub>6</sub> Averaging kernels Figure B.1 shows all kernels for the 5000 measurements presented in this paper, color-coded by site altitude (red=3.8 km; orange=2.2 km; lime=1.2 km; cyan=0.35 km; blue < 0.2 km) as in the main body of the paper. The kernels increase with altitude but with <40% variation over the 0-30 km altitude range. Note that the kernels representing the 3.8 km

(... [3])





Figure <u>G</u>1. HYSPLIT back-trajectories for Nov 10, 2015 (day 314) when the highest ever C<sub>2</sub>H<sub>6</sub> was measured from
 JPL. Yellow oval (upper-left) indicates location of Aliso Canyon Underground Storage Facility. Green ball (lower-

right) denotes JPL, at the convergence of trajectories arriving at 19, 20, & 21 UT. Trajectory calculation used the

718 NAM 12 km resolution, hybrid sigma-pressure meteorology. © OpenStreetMap contributors 2020. Distributed under a

719 Creative Commons BY-SA License.

- 720
- T21 Large C<sub>2</sub>H<sub>6</sub> amounts (3x normal) were observed from JPL on Nov 10 (Day 314), but no enhancement of C<sub>3</sub>H<sub>8</sub>.
- 722 HYSPLIT back-trajectories for this day indicate that the air arriving at JPL at 1000m above ground was from the

723 North-West and had passed over Aliso Canyon USF, confirming that the air over JPL was contaminated by the leak.



Deleted: C

Formatted: Line spacing: 1.5 lines

726	Figure <b>G2</b> . Showing the relationship between CH4 and N2O at JPL in 2014–2017 color-coded by C2H6. Blue points	Deleted: C
727	represent low C <sub>2</sub> H <sub>6</sub> whereas red represents the highest C <sub>2</sub> H <sub>6</sub> . The encircled points represent Nov. 10, 2015, whose	
728	back-trajectory is shown in <u>the</u> previous figure.	
729	Most of the variation in column CH4 and N2O is associated with the stratospheric circulation. Old airmasses from	
730	high latitude are depleted in CH4 and N2O. To remove these effects, and be able to more clearly see changes driven	
731	by the troposphere, $XCH_4$ is plotted versus $XN_2O$ which is similarly affected by stratospheric circulation, but not by	
732	tropospheric emissions. This creates a correlation with the lower-left points representing high-latitude stratospheric	
733	airmasses and the upper right low-latitude airmasses.	
734	The encircled points on Fig. C2 were measured on Nov 10, 2015, when JPL was downwind of the Aliso Canyon	Deleted: C
735	USF leak. The indicate XCH4 enhancements of over 2%, which probably represent a 10+% enhancement in the	
736	PBL with no enhancement above. There is also a general tendency for higher CH4 values when C2H6 is elevated on	
737	other days too, as seen from the dark blue points (low C2H6) being predominantly in the lower right of the figure and	
738	the greener points (higher C <sub>2</sub> H <sub>6</sub> ) being located toward the upper left.	
739		
740	Code Availability	
741	The GFIT code used for the analysis of MkIV spectra is identical to that used by the TCCON project. It is publicly	
742	available under license from the California Institute of Technology for non-commercial use. It can be cloned from:	
743	hg clone https://parkfalls.gps.caltech.edu/tccon/stable/hg/ggg-stable/	
744	after signing the license agreement and being issued a password.	
745		
746	Data Availability	
747	The ground-based MLV data used in this paper can be downloaded from two sites:	Deleted: K
748	https://mark4sun.jpl.nasa.gov/ground.html	
749	ftp://ftp.cpc.ncep.noaa.gov/ndacc/station/barcroft/ames/ftir/	
750	Authors Contributions	
751	Toon, Sung, Blavier for data acquisition. Toon and Yu for data interpretation.	
752	Competing Interests	
753	No competing interests.	
754	Acknowledgements	
755	The authors gratefully acknowledge the NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT	
756	transport and dispersion model and/or READY website (https://www.ready.noaa.gov) used in this publication. We	

760	thank NCEP and GEOS FPIT for their atmospheric analyses. We also acknowledge the NOAA ESRL GMD for
761	distributing in situ data of C <sub>3</sub> H <sub>8</sub> and C <sub>2</sub> H <sub>6</sub> . We thank NASAs Upper Atmosphere Composition Observation (UACO)
762	program for funding support.
763	References
764	Angelbratt, J. et al. Carbon monoxide (CO) and ethane (C2H6) trends from ground-based solar FTIR measurements
765	at six European stations, comparison and sensitivity analysis with the EMEP model. Atmos. Chem. Phys. 11,
766	9253–9269 (2011)
767	Conley, S., G. Franco, I. Faloona, D. R. Blake, J. Peischl, T. B. Ryerson, Methane emissions from the 2015 Aliso
768	Canyon blowout in Los Angeles, Science (2016), 351, (6279), 1317-1320
769	Dalsøren, S.B., Myhre, G., Hodnebrog, Ø. et al. Discrepancy between simulated and observed ethane and propane
770	levels explained by underestimated fossil emissions, Nature Geoscience, 11, 178-184 (2018).
771	https://doi.org/10.1038/s41561-018-0073-0
772	Franco, B. et al. Retrieval of ethane from ground-based FTIR solar spectra using improved spectroscopy: recent
773	burden increase above Jungfraujoch. J. Quant. Spec. Radiat. Trans. 160, 36-49 (2015).
774	Franco, B. et al. Evaluating ethane and methane emissions associated with the development of oil and natural gas
775	extraction in North America. Environ. Res. Lett. 11, 044010 (2016).
776	Harrison, J.J., Allen, N.D.C., and Bernath, P.F., 2010a, Infrared absorption cross sections for ethane (C2H6) in the 3
777	µm region, Journal of Quantitative Spectroscopy and Radiative Transfer, 111, 357-363,
778	DOI: 10.1016/j.jqsrt.2009.09.010
779	Harrison, J.J. and Bernath, P.F., 2010b, Infrared absorption cross sections for propane (C3H8) in the 3 µm
780	region, Journal of Quantitative Spectroscopy and Radiative Transfer, 111, 1282-1288,
781	DOI: 10.1016/j.jqsrt.2009.11.027
782	Helmig, D., Rossabi, S., Hueber, J. et al. Reversal of global atmospheric ethane and propane trends largely due to
783	US oil and natural gas production. Nature Geosci <u>ences</u> , 9, 490–495 (2016). https://doi.org/10.1038/ngeo2721
784	Helmig, D. et al. Climatology and atmospheric chemistry of the non-methane hydrocarbons ethane and propane over
785	the North Atlantic. Elementa 3 (2015).
786	Helmig D., Hueber J., Tans P. (2017), Non-Methane Hydrocarbons from the NOAA ESRL Surface Network, 2004-
787	2016.
788	Howard, Touché, Thomas W. Ferrara, Amy Townsend-Small (2015), Sensor transition failure in the high flow
789	sampler: Implications for methane emission inventories of natural gas infrastructure, Journal of the Air & Waste
790	Management Association, 65:7, 856-862, DOI: 10.1080/10962247.2015.1025925,
791	Irion, F. W., Gunson, M. R., Toon, G. C., Chang, A. Y., Eldering, A., Mahieu, E., Manney, G. L., Michelsen, H. A.,
792	Moyer, E. J., Newchurch, M. J., Osterman, G. B., Rinsland, C. P., Salawitch, R. J., Sen, B., Yung, Y. L., and
793	Zander, R.: Atmospheric Trace Molecule Spectroscopy (ATMOS) Experiment Version 3 data retrievals, Appl.
794	Opt., 41, 6968–6979, 2002

Deleted: ¶
Formatted: Pattern: Clear (White)

 Formatted: Font: (Default) Times New Roman, 10 pt

 Formatted: Font: (Default) Times New Roman, 10 pt

796	Rolph, G., Stein, A., and Stunder, B., (2017). Real-time Environmental Applications and Display sYstem: READY.	
797	Environmental Modelling & Software, 95, 210-228	
798	Rosado-Reyes, C. M., and J. S. Francisco (2007), Atmospheric oxidation pathways of propane and its by-products:	
799	Acetone, acetaldehyde, and propionaldehyde, J. Geophys. Res., 112, D14310, doi:10.1029/2006JD007566.	
800	Sharpe, Steven W. Johnson, Timothy J. Sams, Robert L. Chu, Pamela M. Rhoderick, George C. Johnson, Patricia	
801	A., "Gas-Phase Databases for Quantitative Infrared Spectroscopy", Applied Spectroscopy 58, 1452-1461. (2004)	
802	Stein, A.F., Draxler, R.R, Rolph, G.D., Stunder, B.J.B., Cohen, M.D., and Ngan, F., (2015). NOAA's HYSPLIT	
803	atmospheric transport and dispersion modeling system, Bull. Amer. Meteor. Soc., 96, 2059-2077	
804	Sung, K., G. Toon, A. W. Mantz, and M. A. H. Smith (2013), FTIR measurements of cold C3H8 cross sections at 7-	
805	15 um for Titan atmosphere, Icarus, 226, 1499-1513, doi:10.1016/j.icarus.2013.07.028	
806	Toon, G.C., The JPL MkIV Interferometer, Opt. Photonics News, 2, 19-21, 1991	
807	Toon, G. C., Blavier, JF., Sung, K., Rothman, L. S., and Gordon, I.:, HITRAN spectroscopy evaluation using solar	
808	occultation FTIR spectra, J. Quant. Spectrosc. Ra., 182, 324–336, https://doi.org/10.1016/j.jqsrt.2016.05.021,	
809	2016.	
810	Toon, G. C., Blavier, JF. L., and Sung, K.: Atmospheric carbonyl sulfide (OCS) measured remotely by FTIR solar	
811	absorption spectrometry, Atmos. Chem. Phys., 18, 1923-1944, https://doi.org/10.5194/acp-18-1923-2018,	
812	2018a.	
813	Toon, G. C., Blavier, JF. L., and Sung, K.: Measurements of atmospheric ethene by solar absorption FTIR	
814	spectrometry, Atmos. Chem. Phys., 18, 5075-5088, https://doi.org/10.5194/acp-18-5075-2018, 2018b.	
815	Wunch, D., Toon, G. C., Blavier, JF. L., Washenfelder, R. A., Notholt, J., Connor, B. J., Griffith, D. W. T.,	
816	Sherlock, V., and Wennberg, P. O.: The total carbon column observing network, Philos. T. R. Soc. A, 369,	
817	2087-2112, https://doi.org/10.1098/rsta.2010.0240, 2011.	
818	Touché Howard, Thomas W. Ferrara & Amy Townsend-Small (2015) Sensor transition failure in the high flow	
819	sampler: Implications for methane emission inventories of natural gas infrastructure, Journal of the Air & Waste	
820	Management Association, 65:7, 856-862, DOI: 10.1080/10962247.2015.1025925	
821	The NEED Project. (2017). Propane [pdf]. Retrieved from	
822	http://www.need.org/files/curriculum/infobook/propane.pdf	
823	Urbanski, Shawn P., Wei Min Hao and Stephen Baker, Chemical Composition of Wildland Fire Emissions, Chapter	
824	4, Developments in Environmental Science, Volume 8, 79-107, A. Bytnerowicz, M. Arbaugh, A. Riebau, and	
825	C. Andersen (Editors), ISSN: 1474-8177/DOI:10.1016/S1474-8177(08)00004-1	

Formatted: Font: Times New Roman, 10 pt

Formatted: Font: 10 pt

Page 5: [1] Deleted	Microsoft Office User	2/9/21 9:46:00 AM	
▼			
<b>A</b>			
Page 22: [2] Deleted	<b>Microsoft Office User</b>	3/13/21 3:33:00 PM	
•			
<b>A</b>			
Page 23: [3] Deleted	Microsoft Office User	2/4/21 4:24:00 PM	
X			

I