#### A. Point by Point Response to Reviews

- 3 Dear anonymous reviewers,
- 4 Thank you for your thorough reviews and your support in improve this manuscript.
- 5 6

1 2

#### Response to Reviewer 1

# 7 Line 31. Are you sure that "the summer drawdown for COS is 6 times stronger than for CO2"? The magnitude of the 8 times seems too large to be believable.

- 9 The exact wording of the cited paper Montzka et al. (2007) is:
- 10 'However, while reduced mixing ratios of CO2 during the NH growing season represent a balance between vegetative
- 11 uptake and total respiration (i.e. NEP), the percentage reduction on COS mixing ratio is 4-6 times (5.5+/-1.6) larger during
- 12 June-August (calculated relative to mixing ratios measured at 4-8 km asl) (Figure 6c).
- 13 We reworded the sentence to more accurately correspond to the cited paper (Montzka 2007):
- 14 However, the relative decrease in ambient mixing ratio during summer of the northern hemisphere is 6 times stronger for
- 15 COS than for  $CO_2$ , (Montzka et al., 2007) as COS is generally not emitted by plants like  $CO_2$ , which is released in respiration 16 processes.
- 17 Line 130. "while air was sucked through the chamber to the QCL at a flow rate of 1.5 l min-1". The heights of air
- 18 inlets for the chamber and ambient environment should be noted because remarkable vertical distribution of COS

19 mixing ratio near the ground was observed in this study. If the height of air inlet for the chamber was within the

- 20 canopy of the grass, the COS uptake flux would be largely overestimated, e.g., the COS mixing ratio could drop to
- $21 \quad 134 {\rm ppt} \ {\rm within} \ {\rm the} \ {\rm canopy} \ {\rm in} \ {\rm comparison} \ {\rm with} \ {\rm about} \ 500 {\rm ppt} \ {\rm over} \ {\rm the} \ {\rm canopy}.$
- 22 The intake height was at 0.12 m above the ground and thus within the canopy. The COS concentration inside the chamber
- 23 was thus similar to what the undisturbed soil would experience, which avoids uptake/release being biased high/low when
- 24 COS-enriched air from above the canopy would be used.
- 25 We included this information in the method section.
- 26 The intake height of the ambient as well as the inlet of the chamber air were located at 0.12 m above the ground and thus
- 27 within the canopy height with the exception of right after the cuts (see cutting dates in Section 2.1).
- 28 However, we also have to disagree with the comment on the overestimation of the uptake flux if the intake was within the
- 29 canopy. In order not to bias measurements, the mixing ratios used for chamber flux measurements should be as close to
- 30 reality as possible, which is why air from within the canopy was used. Had we used COS-enriched air from above the
- 31 canopy, any soil uptake would have been overestimated (because the COS gradient across the soil surface is increased),
- 32 while any COS emission would have been underestimated (because the COS gradient across the soil surface is reduced).
- 33 We added this information to the discussion:
- 34 The low COS mixing ratios observed in the lowermost canopy layers just above the soil surface emphasize the importance of
- 35 using air from within the canopy for soil chamber measurements and not COS richer air from above the canopy, which
- 36 would increase the COS gradient and thus increase uptake/decrease emission of COS to/from the soil.
- Line 228. What's the plant available water? Fig. 1 only presents the SWC (%) which is below 38% during almost all
  days, rather than 21 days.
- 39 The SWC in Fig.1 was replaced with the plant available water, which falls below 50 % during 111 days.
- 40 Line 248. "During nighttime (RSW = 0, n = 43), the soils of the grassland acted as a net sink for COS 74.4 % of the
- 41 time" is better replaced by "During nighttime (RSW = 0, n = 43), 74.4 % of the COS emission fluxes were negative,
- 42 implying soils of the grassland acted as a net sink for COS".

43 The sentence was changed as suggested.

44 Line 263. Why did you use both circles and open diamonds for depicting COS soil fluxes? What's the difference 45 between them?

46 We removed the depiction about the open diamonds, which were not present in the plots.

47 Lines 276-278. "Especially after the cuts we observed a strong decline in COS uptake and even times where the 48 grassland turned into a net source for COS with midday means of up to 24.5 pmol m-2s-1 (Fig. 4 b) for up to 8 days 49 after the cut, when the dried litter had already been removed (Fig. 2 a-c)". This sentence is suggested to be replaced 50 by "Especially after the cuts we observed a strong decline in COS uptake ((Fig. 4 b)) and the grassland even turned 51 into a net source for COS in middays (Fig. 2 a-c) with a highest emission flux of 24.5 pmol m-2s-1 in August after the

52 cut.".

53 We replaced the sentence according to your suggestion. To keep the crucial information about the grassland turning into a 54 net source for up to 8 days after the cut, we added an additional sentence to the manuscript:

55 We observed COS emissions for up to 8 days after the cut, when the dried litter had already been removed (Fig. 2 a-c).

56 Lines 280-281. "The cut in October led to a reduction in COS uptake, which was lowest three days after the cut (Fig.

57 2d)". The description seems to be inconsistent with the Fig. 2d.

58 We agree that the lowest COS uptake did occur later than 3 days after the cut. However, this is also related to the overall

59 decline in COS uptake by the grassland at the end of the season. We see no recovery of the COS flux after the last cut. We

60 rephrased this in the manuscript and included more data points to Fig. 2 d):

61 The cut in October led to a reduction in COS uptake, which declined across several days and did not recover, as the end of 62 the season was reached (Fig. 2 d & Fig. 5 b).

63 Lines 297-298. I don't understand the meaning of the sentence. Fig. 4a is the seasonal cycle of CO2, rather than COS.

64 We changed Fig 4a to 4c and added Fig 4a to subsequent sentence dealing with the seasonal response of respiration.

Lines 325-328. I wonder why the COS mixing ratio dropped so large during the nighttime when the COS uptake was
 much less than that during midday.

67 Compared to the constant influx of COS rich air during daytime, due to the increased boundary layer (see line 422), this
68 influx stops during nightime and COS gets depleted within the canopy, even when the COS uptake of the ecosystem is lower

69 than during daytime. The strong input of COS rich air during daytime has also been reported by other studies.

70 We added references to the manuscript (Campbell 2017 & Rastogi 2018).

71 Lines 375- 377. I don't understand the logic of this sentence. Because the chamber enclosed both soil and the residual

72 grass after the cuts, the COS emission under sunlight irradiation might be due to the residual rather than the soil 73 itself, e.g., the photochemical formation of COS from the possible liquid released from the cut grasses (JGR, 109,

74 D13301, doi:10.1029/2003JD004206, 2004; JES, 51 (2017)146-156). If the COS emission was ascribed to soil,

75 the authors are suggested to verify it by using a flow tube method under dark and irradiation conditions.

76 We removed this sentence.

77 Line 413. Why did the lowest COS mixing ratio appear in winter when vegetation COS uptake is relatively low?

78 During winter, no strong emission fluxes are expected to originate from vegetation and soils. The mixing ratios rather

79 depend on the transport of COS enriched air from oceans, which are also highest in summer (see Montzka 2007).

#### 81 Lines 419-421: The above sentences didn't mention the difference in concentrations during day and nighttime.

82 We added the sentence:

83 Even though the COS mixing ratio at the layer closest to the soil were higher during day than during nighttime, the absolute

84 decrease in COS was lower during nighttime due to partial stomatal closure (Kooijmans et al., 2017; Campbell et al., 2017).

85 The absolute difference in concentrations during day and nighttime originate from changes in the height of the planetary 86 boundary layer (PBL).

87

# Lines 421-422. Considering the much stronger COS uptake by the grass in daytime than in nighttime, COS mixing ratio above the canopy should decrease in daytime, rather than nighttime despite of the variation of PBL

90 Several studies (e.g. Rastogi 2018 – Ecosystem fluxes of carbonyl sulfide in an old-growth forest: temporal dynamics and

91 responses to diffuse radiation and heat waves) showed that the PBL is the main influence factor on sub-diurnal variability in

92 COS mixing ratio. The incomplete stomatal closure as well as the soil sink cause the nighttime decrease in mixing ratio as

93 there is no influx of COS rich air from the atmosphere. The stronger daytime drawdown can also be observed in the gradient

94 analysis as the decrease in COS mixing ratio, from to the canopy height down to the soil was higher during daytime (125

95 *ppt) compared to the nighttime decrease (102 ppt).* 

96 This information is already present in the manuscript; see line 325-328 and 419-423.

97 98

## Response to Reviewer 2

99 1. Definition of "LRU on ecosystem scale": note that most LRUs in the literature were derived from branch chamber 100 measurements, and were then used in the relationship between Fcos and Fco2 (Eq.1), with the implication/assumption 101 that LRUs derived from branch chamber measurements are representative of the entire canopy. Here the authors 102 infer the LRU (of the entire canopy) from ecosystem flux measurements. Please clarify this.

103 We added that the LRU was calculated using eddy fluxes without the need to use chambers to the method section:

104 Using the above stated method infers LRU solely on the basis of fluxes on ecosystem scale, whereas other studies typically

105 used branch/leaf chamber measurements (Yang et al., 2018) to determine the relationship between the COS and CO2 uptake 106 rates.

107 2. CO2 observations: IRGA CO2 measurements were used in the analyses. I believe that the QCL also measured 108 CO2. Were those data used somehow? If IRGA CO2 measurements were calibrated to the WMO scale, CO2 should 109 be reported as mole fractions instead of mixing ratios, because the WMO scale (NOAA calibration gases) is reported 110 on mole fractions. The difference between mole fractions and mixing ratios is significant for CO2, and not significant

111 for COS.

119

112 The COS and  $CO_2$  fluxes were calculated using solely the QCL data as stated in section 2.5.2. We followed the processing 113 steps of Gerdel et al. 2017 to retrieve the fluxes using the same filters, which as stated by Gerdel et al. 2017 has the

114 advantage that the influence of the high pass filter on the ecosystem relative uptake (ERU) largely cancels out, if applied on

115 COS as well as  $CO_2$ . The ambient COS and  $CO_2$  concentrations both originated from the QCL data, which puts out mixing

116 ratios. We changed the method section accordingly since neither  $CO_2$  nor  $H_2O$  fluxes of the IRGA were used in the final

117 version of the manuscript. We apologize for the confusion.

### 118 3. What are the reasons for the relatively low enhancements of daily maximum PAR values reaching the soil surface

after the third and the fourth cuts (Figure 1)? These are not consistent with the "incident shortwave radiation

120 reaching the soil surface" in Figure 3e.

121 The data of the PAR reaching the soil surface in Fig 1 originated from a PAR sensor that was likely overgrown by short

122 vascular plants and mosses growing directly at the soil surface at the end of the season. We changed the data from this

123 sensor to the data of Fig 3e, which was calculated using the Beer-Lambert law (see line 151).

4. Fcosmedian turned to positive after the third cutting while remained largely negative after the fourth cutting

125 (Figure 2c&d), given that COS soil fluxes would be both positive. What could explain the difference here?

126 The modelled soil fluxes were always relatively small compared to the ecosystem scale fluxes and shouldn't be the reason for

the difference between fig 2c&d. Also, there is less incoming solar energy at the end of the season, likely also decreasing the
emission strength of the residual litter.

129 We added a sentence containing this to the discussion:

130 We did not observe strong COS emissions after the last cut, as the incoming solar radiation, which we hypothesize to amplify 131 the degradation of sulfur containing compounds of plants, was reduced at the end of the season.

132 5. High-light conditions: what is the definition of high-light conditions? How sensitive is the estimated LRU at high

133 light intensity to the choice of high-light conditions?

134 The parameter "iota" – LRU under high light conditions results from equation 8. The second parameter "kappa" controls

the exponential decrease of LRU when the incoming photosynthetic active radiation (PAR) is decreasing and limiting GPP
but not the COS flux.

$$LRU = \iota e^{\left(\frac{\kappa}{R_{PAR}}\right)}$$

137 While mathematically iota is only obtained at infinitely high PAR, in practice above about 700  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> PAR only 138 insignificant change in the ecosystem relative uptake, reflecting the relationship between the COS and the CO<sub>2</sub> flux, can be

139 observed.

140 We included the definition for high light into the methods part:

141 While mathematically i is only obtained at infinitely high PAR, in practice above about 700 µmol  $m^2 s^{-1}$  PAR (Kooijmans et

142 al., 2019) only insignificant change is reported in other studies (Stimler et al., 2011).

- 143 Other technical comments:
- 144 Line 111: I think it is more likely by a GC-MS than a GC, please double check.
- 145 We changed GC to GC-MS within the revised document.
- 146 L154: The unit of RSW-soil should be Wm-2, and for other places as well.
- 147 We changed this according to the reviewer comment.
- 148 L165: obtain-high resolution ! obtain high-resolution
- 149 We changed this according to the reviewer comment.
- 150 L191: Eq.7 was developed in earlier studies, please refer to the original work.
- 151 We changed this according to the reviewer comment and added (Sandoval-Soto 2005) as reference.
- 152 L198-203: It will read better if these are moved to after L188.
- 153 We changed this according to the reviewer comment.
- 154 L230: It needs a bit more explanation of NDVI, what does it indicate?
- 155 We changed the manuscript accordingly.
- 156 Figure 3 caption. open diamonds?
- 157 We removed the text part about the open diamonds, which are not present in the figure.

#### 158 L312: why is an increase in RECO expected?

159 Even though there is a reduction in plant respiration, the increase in incoming radiation reaching the soil surface leads to

- 160 an increase in soil temperature and consequently soil respiration (see Fig.5a). We added this information to the manuscript:
- 161 While the grassland acted as a net sink for CO2 during periods of high LAI (Fig. 5 6 b), a combination of a decline in GPP
- 162 and an increase in daytime RECO, as more incoming radiation was heating the soil surface, turned it into a net source
- 163 during midday in periods of low LAI (Fig. 56 a).
- 164 L319: should be COS instead of CO2
- 165 We changed this according to the reviewer comment.

166 L433-435: LRU is a normalized ratio, and should not depend on the ambient COS. I do not get the point here. 167 This is not quite right. LRU is calculated in order to normalize for differences in COS (and  $CO_2$ ) concentrations, which affect the fluxes. For the same COS and CO2 flux and the same CO2 concentration, LRU will differ whether the ambient 168 COS concentration is 400 or 500 ppt. This is what we quantified in the linear perturbation analysis and what this sentence 169 170 refers to. 171 L437-439: Please specify which are the exact "those observations". Figure 4 indicates that low COS fluxes took place 172 shortly after the cuttings, which coincides with COS emissions from soils after the cuttings. We clarified this by changing the sentence to: 173 174 For the calculation of LRUs we had to remove the canopy flux data containing COS and/or CO2 emissions observations since these would yield negative values for ERU and LRU (see Eq.8). 175 176 L419-422: It may be worth pointing out that the vertical gradient of COS between the canopy level and below the canopy levels exists throughout the day and night, but that of CO2 does not. 177 178 We added the information to the discussion. 179 We only observed an increase in CO2 mixing ratios, caused by the release of CO2 through respiration processes in the soil, 180 whereas COS mixing ratios further declined down to the soil surface. 181 Response to Reviewer 3 182 Minor comments in general: 183 184 There seems to be a really strong gradient within the grass canopy. Would the really low COS above the soils (100-200 ppt) influence the COS flux? 185 186 Yes, since the exchange across the soil surface is driven by the concentration gradient between the ambient air just above 187 the soil surface and within the soil. We added a sentence containing this information to the discussion: 188 The low COS mixing ratios observed in the lowermost canopy layers just above the soil surface emphasize the importance of 189 using air from within the canopy for soil chamber measurements and not COS richer air from above the canopy, which 190 would increase the COS gradient and thus increase uptake/decrease emission of COS to/from the soil. 191 Out of interest, what does the FCOS/[COS] (COS deposition velocity) look like? 192 We provide the plot in the revised supplement. 193 I also think the concentration discussion (Sections 3.4, Fig 6, 4.3) should come before the flux discussion. It really sets 194 the context to fully appreciate the flux discussion. 195 We agree and moved the parts accordingly. Data needs to be made public before publication! Make sure in the final version that the text in the figures is big 196 197 enough. I was having to zoom in a lot to read things. 198 The data is online now and the font size of the text within the figures was increased. I'm really impressed at how well the FP+ model works for grass (Fig 5b/d). 199 200 Thank you, we were also very happy with the mean diel fluxes resulting from the model. 201 What drives the large change in CO2 variability between day and night? As shown by Wohlfahrt et al. (2005), the large variability of NEE during nighttime conditions is due to the combination of 202 203 low wind speeds and stable stratification which results in highly intermittent CO<sub>2</sub> fluxes compared to well-mixed convective daytime conditions. On a half-hourly basis, fluxes may even be negative (i.e. net uptake of  $CO_2$ ), which is biologically 204 205 impossible, but results from the intermittent nature of the  $CO_2$  transport and is typically compensated for by large emission fluxes in a subsequent averaging period. As recommended by Wohlfahrt et al. (2005), CO<sub>2</sub> fluxes were filtered for u\*, but not 206 207 for the sign of the fluxes in order not to bias nighttime fluxes towards too large  $CO_2$  emission.

208 We added this reference and information to the manuscript.

#### #Has the data been filtered for u\*? Has any of this large variability been taken into account in the Reco vs temp 209 calculation for GPP uncertainty (something to think about in future if not?). 210

The data has unintentionally not been filtered for u<sup>\*</sup>. We determined the threshold at  $\sim 0.2 \text{ m s}^{-1}$  for CO<sub>2</sub> and used the same 211

value for COS. After reanalyzing the data, we observed only minor changes and no changes in the overall patterns. Text and 212

figures were adapted accordingly. We attached all plots before and after the correction at the end of this document. During 213

the reanalysis we were also able to recover more data from immediately after the first cut, which slightly increased LRU and 214

- 215 ERU during this phase in Fig. 7a.
- There is a little repetition with the Results and Discussion being separate. I wouldn't object if the authors decided to 216
- 217 combine both and tightened the text up. But obviously that's just a suggestion.
- We thank reviewer 3 for the advice but prefer to keep the sections separated. 218
- 219 We removed several redundancies.
- Minor comments by line number: 220
- 221 14: soil flux
- 222 We changed this according to the reviewer comment.
- 223 31: do you mean relative uptake? COS is in ppt vs CO2 in ppm
- Yes, we reworded the sentence to more accurately correspond to the cited paper (Montzka 2007): 224
- 225 However, the relative decrease in ambient mixing ratio during summer of the northern hemisphere is 6 times stronger for
- 226 COS than for CO2, (Montzka et al., 2007) as COS is generally not emitted by plants like CO2, which is released in
- 227 respiration processes.
- 228 38: Extra bracket
- 229 We added a comma and removed the bracket.
- 230 86: What kind of fertilizer (dairy? beef? pig?)? And when was it fertilized previously? Before the winter?
- 231 The grassland is fertilized with solid manure and cattle slurry (see Hörtnagl et al. 2018) once a year at the end of the 232 growing season in October. We added the information to the manuscript:
- 233 Each year, the field site was fertilized with solid manure and cattle slurry (Hörtnagl et al., 2018) at the end of the season 234 (07.10. in 2015).

#### 235 140: Ambient COS from what height? There is a massive COS gradient so this will be important.

- 236 The intake height was at 0.12m above the ground and thus within the canopy with the exception of measurements taken just
- 237 after the cuts. This information is now included in the method section:
- 238 The intake height of the ambient as well as the inlet of the chamber air were located at 0.12 m above the ground and thus within the canopy height with the exception of measurements right after the cuts (see cutting dates in Section 2.1). 239
- 160: I think this needs more explanation. What does an OBB represent? Is that good? Not good? If you aren't going 240
- into enough detail for readers to evaluate the model, then cut it. It's kind of hanging there with not enough info. And 241
- 242 most of the packages mentioned will represent some mathematical approach to data analysis. Since packages come
- 243 and go, it would be really helpful to have a sentence or two about what these packages actually represent.
- The OOB score can be interpreted as a pseudo-R2 and is widely used in random forest analyses (regression and 244
- classification), especially in the absence of a proper test dataset. It uses the data not seen by the trees (random forest uses 245 246 bootstrapping) as a test dataset. We added this information to the methods section.
- 168: What heights along the tower were the gradients sampled from? How often were they sampled vs eddy flux 247 248
- sampling?
- 249 The information was already present in the methods section. See 2.3 and 2.3.1

- 250 173: Was the eddy flux data filtered for insufficient turbulence? If so, what u\* filter was applied? How was the u\*
- threshold quantified? A plot of the FCOS and FCO2 vs u\* would be helpful here to understand the micro met dynamics for the site.
- 253 The u\* threshold was determined by running the change point detection algorithm of Barr et al (2013) on nighttime NEE.
- The  $u^*$  for the  $CO_2$  flux (~0.2 m s<sup>-1</sup>) was then applied for COS. We also tried to determine the  $u^*$  threshold for COS, but a satisfying change point couldn't be determined.
- 256 We noticed that the eddy flux data was unintentionally not correctly filtered for u\* in the plots (which almost exclusively has
- 257 only an effect during the night). The data in the plots and the corresponding values in the text have been updated.
- 258 We added the plot of the  $FCO_2$  vs  $u^*$  to the supplement.
- 329: What does the [CO2] drop down to? Is there a relationship between u\*/turbulence and the d[COS] and d[CO2]?
  That would be an interesting figure to see.
- 261 The  $CO_2$  mixing ratio drops down to 339 ppm at 0.1m above ground at 10 a.m. We added a plot containing the u\* values
- and the differences of the  $CO_2$  and COS mixing ratios between canopy level (0.4m) and 0.02 m for COS and 0,1m for  $CO_2$  to
- the supplement. The two lowest measurement heights were excluded for  $CO_2$  since there the  $CO_2$  mixing ratio increased due to the soil respiration.
- 422: How long does the morning increase in COS last for? Do you start to see a decrease in COS as the daytime uptake influences the air in the valley? Other sites have also seen this morning peak in COS. Maybe include a
- 267 reference to those here. (e.g. Redwoods, Harvard Forest, etc)
- 268 We observed a steep morning increase in COS mixing ratios until about 11 a.m.. We included include this plot in the 269 supplement and added the requested information to the discussion.
- 270 While the PBL is shallow during nighttime and the COS mixing ratio decreases due to the sink strength of the grassland, at
- 271 the onset of the day, the PBL layer height increases quickly and COS rich air is transported down to the ecosystem (see Fig.
- 272 S12) (Campbell et al., 2017). A similar steep increase until midday has also been observed by Rastogi et al. (2018).

274 Updated figures:

275 New:







285 New:









## B. List of relevant changes

- The correct u\* filter is now applied and all values in the document have been changed accordingly
- During the reanalysis we were also able to recover more data from immediately after the first cut, which slightly
   increased LRU and ERU during this phase (Fig. 7a)
- The section about the COS and CO2 mixing ratios is now placed before the flux sections in the result as well as the
   discussion section
- 302

#### C. Tracked document

303

304

# Seasonal dynamics of the COS and CO<sub>2</sub> exchange of a managed temperate grassland

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310 Abstract. Gross primary productivity (GPP), the CO<sub>2</sub> uptake by means of photosynthesis, cannot be measured directly on 311 ecosystem scale, but has to be inferred from proxies or models. One newly emerged proxy is the trace gas carbonyl sulfide 312 (COS). COS diffuses into plant leaves in a fashion very similar to  $CO_2$ , but is generally not emitted by plants. Laboratory 313 studies on leaf level gas exchange have shown promising correlations between the leaf relative uptake (LRU) of COS to CO<sub>2</sub> 314 under controlled conditions. However, in situ measurements including daily to seasonal environmental changes are required, 315 to test the applicability of COS as a tracer for GPP at larger temporal scales. To this end, we conducted concurrent 316 ecosystem scale CO<sub>2</sub> and COS flux measurements above an agriculturally managed temperate mountain grassland. We also 317 determined the magnitude and variability of the soil COS exchange, which can affect the LRU on ecosystem level. The 318 cutting and removal of the grass at the site had a major influence on the soil flux as well as the total exchange of COS. The 319 grassland acted as a major sink for CO<sub>2</sub> and COS during periods of high leaf area. The sink strength decreased after the cuts 320 and the grassland turned into a net source for CO<sub>2</sub> and COS on ecosystem level. The soil acted as a small sink for COS when 321 the canopy was undisturbed, but also turned into a source after the cuts, which we linked to higher incident radiation hitting 322 the soil surface. However, the soil contribution was not large enough to explain the COS emission on ecosystem level, 323 hinting to an unknown COS source possibly related to dead plant matter degradation. Over the course of the season, we 324 observed a concurrent decrease of CO<sub>2</sub> and COS uptake on ecosystem level. With the exception of the short periods after the cuts, the LRU under high light conditions was rather stable and indicates a high correlation between the COS flux and GPP 325 326 across the growing season.

327

#### 328 1 Introduction

329 Carbonyl sulfide (COS) is the most abundant sulfur-containing gas in the atmosphere with tropospheric mixing ratios of 330 ~500 ppt. Within the atmosphere, COS acts as a greenhouse gas with a 724 times higher direct radiative forcing efficiency as CO2 (Brühl et al., 2012). After reaching the stratosphere, it reacts to sulfur aerosols via oxidation and photolysis, hence 331 contributing to the backscattering of solar radiation and having a cooling effect on Earth's atmosphere (Krysztofiak et al., 332 333 2015; Whelan et al., 2018). The intra-seasonal atmospheric COS mixing ratio follows the pattern of CO<sub>2</sub> as terrestrial vegetation acts as the largest known sink for both species (Montzka et al., 2007; Whelan et al., 2018; Le Quere et al., 2018). 334 However, the relative decrease in ambient mixing ratio during summer of the northern hemispherethe summer drawdown for 335  $\frac{\text{COS}}{\text{COS}}$  is 6 times stronger for  $\frac{\text{COS}}{\text{COS}}$  than for  $\text{CO}_2$  (Montzka et al., 2007) as COS is generally not emitted by plants like  $\text{CO}_2$ , 336

337 which is released in respiration processes.

338 The uptake of COS by plants is mostly mediated by the enzyme carbonic anhydrase (CA), but also photolytic enzymes like

- 339 Ribulose-1,5-bisphosphate-carboxylase/-oxygenase (Rubisco) (Lorimer and Pierce, 1989). This in turn means that COS and
- 340 CO<sub>2</sub> share a similar pathway into leaves through the boundary layer, the stomata and the cytosol, up to their reaction sites.

Compared to  $CO_2$ , COS is processed in a one-way reaction to  $H_2S$  and  $CO_2$  (Protoschill-Krebs and Kesselmeier, 1992;Notni et al., 2007) and therefore not released by plants, (with the exception of severely stressed plants (Bloem et al., 2012;Gimeno et al., 2017)). That makes COS an interesting tracer for estimating the stomatal conductance and the gross uptake of  $CO_2$ , referred to as gross primary production (GPP), on ecosystem level (Asaf et al., 2013;Kooijmans et al., 2017;Kooijmans et al., 2019). However, to estimate GPP using COS, the relative uptake of COS to GPP deposition velocities (LRU) must be known beforehand (see Eq.1), so that GPP can be estimated on the basis of the COS flux.

$$347 \quad LRU = \frac{\frac{r_{COS}}{\chi_{COS}}}{\frac{F_{CO2}}{\chi_{CO2}}}$$
(Eq.1)

 $F_{COS}$  is the COS leaf flux (pmol m<sup>-2</sup> s<sup>-1</sup>),  $F_{CO2}$  is the gross CO<sub>2</sub> uptake on leaf level (µmol m<sup>-2</sup> s<sup>-1</sup>) and  $\chi_{COS}$  and  $\chi_{CO2}$  are the 348 ambient COS and CO2 mixing ratios in ppt and ppm, respectively. Leaf level studies for C3 plants have estimated the LRU to 349 be around 1.7 with the 95% confidence interval between 0.7 and 6.2 (Whelan et al., 2018;Seibt et al., 2010;Sandoval-Soto et 350 351 al., 2005). The large spread of the LRU most likely originates from differences between plant species, for example, leaf structure and plant metabolism (Wohlfahrt et al., 2012;Seibt et al., 2010), which questions the applicability of the concept of 352 LRU in real-world ecosystems under naturally varying environmental conditions. It is also known that the LRU is just stable 353 under high light conditions, since the uptake of CO<sub>2</sub> by means of photosynthesis is a light driven process, while CA is able to 354 process COS independently of light conditions (Maseyk et al., 2014; Yang et al., 2018; Stimler et al., 2011). Any model of 355 356 LRU should therefore reflect diurnal changes in light conditions. Kooijmans et al. (2019) recently discovered that the vapor pressure deficit (VPD) appears to have a stronger control on F<sub>CO2</sub>, than on F<sub>CO2</sub>, in an evergreen needle forest. If generally 357 true, this would add further variability to the LRU and complicating the application of COS to estimate GPP. Besides inter-358 specific differences in LRU, the question remains unanswered if the LRU is also susceptible to seasonal changes of 359 360 ecosystems for example, changes in species composition or phenology, which would further complicate the application of COS in carbon cycle research. Maseyk et al. (2014) observed COS emissions on ecosystem scale over a winter wheat field 361 362 going into senescence, indicating that potentially strong sources of COS could distort LRU.

Since CA and other enzymes known to emit or take up COS are also present in microorganisms (Ogawa et al., 2013;Seefeldt 363 et al., 1995;Ensign, 1995;Smeulders et al., 2013;Whelan et al., 2018), recent studies have also quantified the contribution of 364 soils to the COS ecosystem flux (Kooijmans et al., 2017;Spielmann et al., 2019;Maseyk et al., 2014). COS soil fluxes could 365 366 modify the LRU on ecosystem level and hence inferred GPP, if they are substantial compared to COS canopy fluxes. Similar to the ecosystem fluxes, the soil fluxes could not only be prone to diurnal, but also seasonal changes, depending on the 367 substrate availability, environmental conditions (e.g. soil temperature and moisture), substrate quality and quantity, and 368 changes in composition of the microbial communities (Kitz et al., 2019;Meredith et al., 2019). Recent studies have also 369 linked COS soil emissions to abiotic processes dependent on light and/or temperature (Whelan and Rhew, 2015;Kitz et al., 370 371 2019;Meredith et al., 2018).

372 The goal of our study was to provide new insights into the seasonal variability of COS fluxes on ecosystem, soil and canopy 373 level. To this end, we conducted a 6-month campaign on a managed temperate mountain grassland, measuring ecosystem as 374 well as soil COS fluxes. Since the grassland was cut four times during the campaign, we were able to observe multiple 375 growing cycles and investigate the diel and seasonal changes of the COS fluxes and the LRU in this highly dynamic 376 ecosystem. We hypothesize that (H1) the grassland, given its large CO<sub>2</sub> uptake capacity (Wohlfahrt et al. 2008), is a major 377 sink for COS and that the sink strength decreases over the course of the season, (H2) the drying of the cut grass leads to a 378 release of COS, (H3) the LRU will change after the cuts, due to stressed plants and drying plant parts in the field, but is 379 otherwise stable,- (H4) the cuts turn the soil into a COS source, due to the larger amount of light reaching the soil surface 380 (Kitz et al., 2017), but once a reasonably high leaf area index (LAI) has developed, COS is taken up by soil.

#### 381 2 Methods

#### 382 2.1 Study site and period

383 The study was conducted at an intensively managed mountain grassland in the municipal territory of Neustift (Austria) in Stubai valley (FLUXNET ID: AT-Neu; doi: 10.18140/FLX/1440121). The grassland is situated at an elevation of 970 m a.s.l. 384 in the middle of the flat valley bottom. The soil was classified as Fluvisol with an estimated depth of 1 m with the majority 385 of roots located within the first 10 cm. Measurements were conducted between 01.05.2015 and 31.10.2015 (183 days). The 386 vegetation was described as Pastincao-Arrhenatheretum and consisted mainly of Dactylis glomerata, Festuca pratensis, 387 Alopecurus pratensis, Trisetum flavescens, Ranunculus acris, Taraxacum offcinale, Trifolium repens, Trifolium pratense, 388 389 and Carum carvi (Kitz et al., 2017). During the campaign, the grassland was cut four times (02.06./07.07./21.08./01.10.2015) and the biomass left to dry on the field for up to one day, before being removed as silage. Each year, tThe field site was 390

## 391 fertilized with organic solid manure and cattle slurry (Hörtnagl et al., 2018) at the end of the season (07.10- in 2015).

#### 392 2.2 Leaf area index

1 /

The LAI was estimated from assessments of the average canopy height, which were related to destructive LAI measurements, using the following sigmoid function:

395 
$$LAI = \frac{1}{(1 + \exp(-(a_1DOY + a_2)))(b_1 - b_2)}$$
 (Eq.2)

where DOY is the day of the year and a1, a2, b1 and b2 are factors that were optimized for each growing period, for example, before the first cut, between cuts and after fourth cut (Wohlfahrt et al., 2008). Additionally, biomass samples were taken at 15 occasions, to assist with the LAI calculation.

#### 399 2.3 Mixing ratio measurements

The CO<sub>2</sub> ( $\chi_{CO2}$ ) and COS ( $\chi_{COS}$ ) mixing ratios were measured using a Quantum Cascade Laser (QCL) Mini Monitor (Aerodyne Research, Billerica, MA, USA) at a wavenumber of ca. 2056 cm<sup>-1</sup> and at a frequency of 10 Hz. To minimize the effect of air temperature (T<sub>air</sub>) changes on the instrument, we placed it in an insulated box which in turn was located in a climate controlled instrument hut (30°C). The cooling of the laser was achieved by a chiller (ThermoCube 400, Solid State Cooling Systems, Wappinger Falls, NY, USA).

We used ¼ inch Teflon™ tubing, stainless steel fittings (SWAGELOK, Solon, OH, USA and FITOK, Offenbach, HE, 405 Germany), Teflon Filters (Savilex, EdenPrarie, MN, USA) as well as COS-inert valves (Parker-Hannafin, Cleveland, OH, 406 USA) to ensure that only materials known not to interact with COS were used for the measurement and calibration airflow. 407 408 The H<sub>2</sub>O and CO<sub>2</sub> mixing ratios (7120.& 7002.) were measured by a closed path infrared gas analyzer (IRGA) (Licor 6262, 409 LICOR Biosciences, Lincoln, NE, USA). Since the data of the QCL, and the sonic anemometer and the IRGA-were saved 410 on two separate PCs, a network time protocol software (NTP, Meinberg, NI, Germany) was used to keep the time on both 411 devices synchronized. We corrected known  $\chi_{COS}$  drift issues of the QCL (Kooijmans et al., 2016) by doing half hourly calibrations for 1 min with a gas of known  $\chi_{COS}$ . The gas cylinders (working standards) used for the calibrations were either 412 pressurized air (UN 1002) or nitrogen (UN 1066), which were cross-compared (when working standard cylinders were full 413 414 and close to empty) to an Aculife-treated aluminum pressurized air cylinder obtained from the National Oceanic & 415 Atmospheric Administration (NOAA). The latter was analyzed by the central calibration laboratory of NOAA for its  $\chi_{COS}$ using gas chromatography with mass spectrometric detection (GC-MS) on 06.04.2015. We then linearly interpolated 416 417 between the offsets of the half hourly calibrations and used the retrieved values to correct the high frequency COS data. Due 418 to issues with the scale gas cylinder, no absolute concentrations were available before the 16.06.first cut The COS mixing 419 ratios were extrapolated to the 1<sup>st</sup> cut to increase the amount of available data for the first post cut and therefore no LRU was

#### 420 ealculated for this period. This was done on the basis of the measured CO<sub>2</sub> mixing ratios and the mean half hourly ratio of

421 the ambient CO<sub>2</sub> to COS mixing ratios between the 16. and the 18.6..

#### 422 2.3.1 Mixing ratio measurements within the canopy

In order to investigate the  $\chi_{COS}$  within the canopy, we used a multiplexer and 8 <sup>1</sup>/<sub>4</sub> inch Teflon<sup>TM</sup> tubes to measure the  $\chi_{COS}$  at 8 heights within and above the canopy i.e. at 2, 5, 10, 20, 30, 40, 50 & 250 cm height above ground with a tube length of 15 m for each height. The upper two intakes were located at the eddy covariance measurement and canopy height, respectively. Each height was measured for 1 min at 1 Hz and 2 1 min<sup>-1</sup>, while the other lines were each flushed at 2 1 min<sup>-1</sup>. The  $\chi_{COS}$  drift was also corrected by doing half hourly calibrations (see section 2.3).

428

#### 429 2.4 COS soil fluxes

#### 430 2.4.1 Soil chamber setup

To quantify soil COS fluxes, we installed four stainless steel (SAE grade: 316L) rings 5 cm into the soil. They remained on site for 112 days (10.06.2015 – 30.09.2015). Two additional rings were installed on the 31.08.2015 and the 02.10.2015 to examine any long-term effects of the ring placement and to replace the original rings for the measurements in September and October. The aboveground biomass within each ring was removed at the day of installation and again at least one day prior to each measurement day. The roots within as well as the vegetation surrounding the rings were not removed and natural litter was left in place. At days without measurements the soil within the rings was covered by fleece to prevent it from drying out.

To measure the soil fluxes, a transparent fused silica-glass chamber (Kitz et al., 2017) was placed into the water filled 438 439 channel of the steel rings, while air was sucked through the chamber to the QCL at a flow rate of 1.5 l min<sup>-1</sup>. The chamber 440  $\chi_{COS}$  was then compared with the ambient  $\chi_{COS}$  above the chamber, using a second inlet to which we switched before the 441 chamber measurement and after reaching stable readings inside the chamber. The intake height of the ambient as well as the 442 inlet of the chamber air were located at 0.12 m above the ground and thus within the canopy height with the exception of 443 measurements right after the cuts (see cutting dates in Section 2.1). Overall, 243 chamber measurements were conducted 444 over the course of the campaign including day and nighttime measurements. Additional manual measurements included a 445 hand-held sensor (WET-2, Delta-T Devices, Cambridge, England) to measure soil water content (SWC) and soil temperature 446 (T<sub>soil</sub>) at a soil depth of 5 cm simultaneously with the soil chamber measurements next to the rings.

#### 447 2.4.1 COS soil flux calculation

448 The COS soil flux was calculated using the following equation:

449 
$$F = \frac{q(\chi_{cos2} - \chi_{cos1})}{4}$$

. .

(Eq.3)

where F is the COS soil flux (pmol m<sup>-2</sup> s<sup>-1</sup>), q denotes the flowrate in (mol s<sup>-1</sup>),  $\chi_{COS2}$  and  $\chi_{COS1}$  are the chamber and ambient  $\chi_{COS}$  in ppt, respectively and A the soil surface area (0.032 m<sup>2</sup>) covered by the chamber. A more detailed description can be found in Kitz et al. (2017).

#### 453 2.4.2 COS soil exchange modelling

Due to the removal of the aboveground biomass and the consequent higher shortwave radiation reaching the soil surface in the chambers, compared to the soil below the canopy, we simulated the soil COS exchange for natural conditions. The soil flux was modelled using our measured soil fluxes and additionally retrieved soil and meteorological data -  $T_{soil}$ , soil water content (SWC) at 5 cm depth next to the chambers and incident shortwave radiation reaching the soil surface ( $R_{SW-soil}$ ) - as input for a random forest regression model (Liaw and Wiener, 2002). The soil fluxes were modelled on half hourly basis for the whole duration of the measurement campaign to calculate the COS canopy fluxes from the difference of the COS ecosystem and soil fluxes. To this end we used the scikit-learn (sklearn Ver. 0.19.1) package, the pandas library and the Python Software Distribution Anaconda (Ver. 5.2.0) in the command shell Ipython (Ver. 6.4.0) based on the Programming language Python (Ver. 3.3.5). We used the Beer-Lambert law to model  $R_{SW_soil}$  under undisturbed conditions as the aboveground vegetation was removed to measure the COS exchange of bare soil:

 $464 \quad R_{SW-soil} = R_{SW} \exp(-K \, LAI)$ 

(Eq.4)

465 where  $R_{SW-soil}$  (Wm<sup>-2</sup>s<sup>-4</sup>) is the shortwave radiation (SW) reaching the soil surface,  $R_{SW}$  is the incoming SW radiation 466 reaching the top of the canopy, LAI is the plant area index (Eq. 2) and K is the canopy extinction coefficient assuming a 467 spherical leaf inclination distribution (Wohlfahrt et al., 2001), which was calculated using the following equation:

468 
$$K = \frac{1}{2\cos(\psi)}$$
 (Eq.5)  
469 where  $\psi$  is the zenith angle of the sun in radians.

470

471 A random forest with 1000 trees was grown which resulted in an out of bag (OOB) score of (0.82). <u>The OOB score can be</u> 472 interpreted as a pseudo-R2 and is widely used in random forest analyses (regression and classification), especially in the 473 absence of a proper test dataset. It uses the data not seen by the trees (random forest uses bootstrapping) as a test dataset. The 474 optimal input parameters, including maximum tree depth, were determined with the function GridSearchCV from the sklearn 475 package.

#### 476 2.5 Ecosystem fluxes

#### 477 2.5.1 Setup for ecosystem fluxes

The COS, CO<sub>2</sub> and H<sub>2</sub>O ecosystem fluxes were obtained using the eddy covariance method (Aubinet et al., 1999;Baldocchi, 2014). We used a 3-axis sonic anemometer (Gill R3IA, Gill Instruments Limited, Lymington, UK) to obtain\_<u>high-high-</u> resolution data of the 3 wind components. The intake of the tube for the eddy covariance measurements was installed in close proximity to the sonic anemometer and insulated as well as heated above  $T_{air}$  to prevent condensation within the tube. The air was sucked to the QCL at a flowrate of 7 l min<sup>-1</sup> using a Vacuum Pump (Agilent Technologies, CA, USA).

#### 483 2.5.2 Ecosystem flux calculation

In a first step we used a self-developed software to determine the time lag, introduced by the separation of tube intake and the sonic anemometer and the tube length, between the QCL and sonic dataset (Hortnagl et al., 2010). The data were then processed using the software EdiRe (University of Edinburgh, UK) and Matlab2019a (MathWorks, MA, USA). We used the laser drift corrected  $\chi_{COS}$  data and linear detrending to process the data before following the procedure to correct for sensor response, tube attenuation, path averaging and sensor separation following Gerdel et al. (2017). The random flux uncertainty was calculated following Langford et al. (2015).

490 We estimated the COS canopy flux from the difference between the measured COS ecosystem and the modelled COS soil 491 flux.

#### 492 2.5.3 Flux partitioning and leaf relative uptake

The GPP on ecosystem level was determined using the FP+ model put forward by Spielmann et al. (2019). The model estimates the GPP on the basis of nighttime net ecosystem exchange (NEE) measurements of  $CO_2$  that are assumed to provide the temperature response of the ecosystem respiration (RECO) as well as a light dependency curve to estimate GPP based on the daytime NEE (Lasslop et al., 2010):

497	$NEE = \frac{\alpha\beta R_{PAR}}{\alpha R_{PAP+R}} + rb \ e^{E_0(\frac{1}{T_{ref} - T_0} - \frac{1}{T_{atr} - T_0})} $ (Eq.6)
498	where $\alpha$ denotes the canopy light utilization efficiency ( $\mu$ mol CO <sub>2</sub> $\mu$ mol <sup>-1</sup> photons), $\beta$ the maximum CO <sub>2</sub> uptake rate of the
499	canopy at light saturation ( $\mu$ mol CO <sub>2</sub> m <sup>-2</sup> s <sup>-1</sup> ), R <sub>PAR</sub> the incoming photosynthetic active radiation ( $\mu$ mol m <sup>-2</sup> s <sup>-1</sup> ), rb the
500	ecosystem base respiration (µmol m <sup>-2</sup> s <sup>-1</sup> ) at the reference temperature $T_{Ref}$ (°C), which is set to 15°C, $T_{air}$ (°C) refers to the
501	air temperature and $E_0$ (°C) to the temperature sensitivity of RECO. $T_0$ was kept constant at -46.02°C. We did not use the
502	VPD modifier of beta put forward by Lasslop et al. (2010) as its value could not be estimated with confidence. We
503	determined the parameter $E_0$ by using nighttime data minimizing the root squared mean error. For the determination of the
504	remaining five unknown model parameters of the two flux partitioning models we used DREAM, a multi-chain Markov
505	Chain Monte Carlo algorithm (for more detail see Spielmann et al. (2019)). We calculated the parameters for ~15 day
506	windows but adjusted them to not overlap with a cut of the grassland.
507	The ecosystem relative uptake (ERU) was calculated using Eq. 1 substituting the GPP with the NEE and using the COS
508	ecosystem flux for F <sub>COS</sub> .
509	The FP+ model by Spielmann et al. (2019) extends the daytime FP (Eq.6) to also estimate the COS ecosystem fluxes by
510	linking the GPP resulting from the first part on the right-hand side of Eq.6 with the COS exchange through:
	GPP LRU CPP LRU COS
511	$F_{cosmodel} = \frac{\chi_{cos}}{\chi_{cos}} \frac{1}{\chi_{cos}} \frac{1}{\chi_{cos}} $ (Eq.7)
512	developed by Sandoval-Soto et al. (2005), where $F_{COSmodel}$ is the modelled COS flux (pmol m <sup>-2</sup> s <sup>-1</sup> ), $\gamma_{COS}$ (ppt) and $\gamma_{CO2}$ (ppm)
513	are the measured ambient mixing ratios of COS and CO <sub>2</sub> respectively and LRU (-) is the leaf relative uptake rate:
~ 1.4	$\left(\frac{\kappa}{2\kappa_{+}}\right)$
514	$LRU = l e^{RpAR} $ (Eq.8)
515	where t (-) corresponds to the LRU at high light intensity and the parameter k (µmol m s) governs the increase of LRU at
516	low light conditions. While mathematically i is only obtained at infinitely high PAR, in practice above about 700 µmol m <sup>2</sup> s
517	<u>PAR</u> (Kooijmans et al., 2019) only insignificant change is reported in other studies (Stimler et al., 2011). The light
518	dependency of LRU originates from the fact that the COS uptake by the enzyme CA is light-independent, while the $CO_2$
519	uptake by Rubisco depends on solar radiation absorbed by leaf chlorophyll (Whelan et al., 2018;Kooijmans et al.,
520	2019;Wohlfahrt et al., 2012).
521	-The method stated above infers LRU solely on the basis of ecosystem scale fluxes, whereas other studies typically use
522	branch/leaf chamber measurements (Yang et al., 2018) to determine the relationship between the COS and CO <sub>2</sub> uptake
523	<u>rates.</u> We determined the parameter $E_0$ by using nighttime data minimizing the root squared mean error. For the determination
524	of the remaining five unknown model parameters of the two flux partitioning models we used DREAM, a multi chain
525	Markov Chain Monte Carlo algorithm (for more detail see Spielmann et al. (2019)). We calculated the parameters for 15
526	day windows but adjusted them to not overlap with a cut of the grassland.
527	The coosystem relative uptake (ERU) was calculated using Eq. 1 substituting the GPP with the NEE and using the COS
528	ecosystem flux for F <sub>COS</sub> .
529	2.5.4 Linear perturbation analysis
/	

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530 The relative contribution of the parameters GPP,  $F_{COSmodel}$ ,  $\chi_{CO2}$  and  $\chi_{COS}$  that drive  $\iota$  (Eq. 7) were estimated through a linear 531 perturbation analysis (Stoy et al., 2006).

532 The changes in  $\iota$  ( $\delta\iota$ ) between the target and the reference window (before the 2<sup>nd</sup> cut, i.e. 18.06.2015-07.07.2015) are 533 considered the total derivative of Eq. 7 and can be represented by a multivariate Taylors's expansion where the higher-order 534 terms are neglected in this first-order analysis:

535 
$$\delta \iota = \frac{\partial \iota}{\partial F_{\text{COSmod}}} dF_{\text{COSmod}} + \frac{\partial \iota}{\partial \chi_{\text{COS}}} d\chi_{\text{COS}} + \frac{\partial \iota}{\partial \text{GPP}} dGPP + \frac{\partial \iota}{\partial \chi_{\text{CO2}}} d\chi_{\text{CO2}}$$
(Eq.9)

536 The relative contributions of the parameters were determined by computing the partial derivatives of Eq. 7.

537	$\frac{\partial \iota}{\partial F_{cOSmod}} = \frac{\chi_{CO2}}{\chi_{COS}GPP}$	(Eq.10)
538	$\frac{\partial \iota}{\partial \chi_{COS}} = \frac{-\chi_{CO2} F_{COSmod}}{\chi_{COS}^2 GPP}$	(Eq.11)
539	$\frac{\partial \iota}{\partial GPP} = \frac{\chi_{CO2} F_{COSmod}}{\chi_{COS} GPP^2}$	(Eq.12)
540	$\frac{\partial \iota}{\partial \chi_{CO2}} = \frac{F_{COSmod}}{\chi_{COS}GPP}$	(Eq.13)

541

#### 542 2.6 Ancillary data

Supporting meteorological measurements included  $T_{air}$  (RFT-2, UMS, Munich, GER),  $T_{soil}$  (TCAV, Campbell Scientific, Logan, UT, USA), SWC (ML2x, Delta-T Devices, Cambridge, UK), incident solar radiation (CNR-1, Klipp and Zonen, Delft, NLD), incident photosynthetic active radiation (PAR) (BF2H, Delta-T Devices Ltd, Cambridge, UK) and the Normalized Difference Vegetation Index (NDVI) sensor (SRS-NDVI, Meter, Pullman, WA, USA). The data were recorded throughout the whole season as 1 min values and stored as half-hourly means and standard deviations.

#### 548 3 Results

#### 549 3.1 Environmental conditions

550 Air temperature ranged between -2 °C and 33 °C with a mean of 13 °C during the study period from 15<sup>th</sup> of May to first of 551 November (Fig. 1). While the majority of precipitation (total 360.5 mm) fell as rain, we observed an exceptionally late snow event on the 20<sup>th</sup> of May, which did not melt for almost two days (Fig. 1). Although the VPD reached values of above 2 kPa 552 during 25 days, and plant available water dropped below 38-50 % on 21-111 days during the campaign (Fig. 1), we did not 553 554 observe any relationship with COS (see Fig S1-S2). Due to the removal of the aboveground biomass, the cuts reduced LAI. 555 They also reduced the normalized difference vegetation index (NDVI) (Fig. 1), which is a measure of canopy greenness 556 (Tucker, 1979), which The NDVI further decreased in the subsequent days as a consequence of dying plant parts remaining at 557 the field site (Fig 2 panels a-c). This can also be observed in the webcam photos (Photo S1-S3).



Figure 1. Seasonal cycle of ancillary variables. Daily minimum, maximum and median (a) air and (b) soil temperatures (°C) indicated by the lower and upper end of the bars and the white circle, respectively. (c) Daily maximum incident photosynthetic active radiations. Shortwave radiation (µmol-W m<sup>2</sup>-s<sup>-1</sup>) reaching the top of the canopy (black squares) and reaching the soil surface (white circles). (d) Daily minimum, maximum and median vapor pressure deficit (kPA) indicated by the lower and upper end of the bars and the white 563 circle, respectively. (e) Soil water contentPlant available water (%) depicted by black squares and cumulative precipitation (mm) depicted by open circles. (f) Modelled leaf area index (black lines), measured LAI (grey squares) and normalized difference vegetation index (open circles).



566

Figure 2: The response of the daily midday medians of NDVI (yellow circles), COS (blue circles) and CO<sub>2</sub> (red circles) ecosystem fluxes
 around the 4 cutting events (a-d) of the grassland. The errorbars depict the respective median absolute deviations. The cuts are marked by a
 red dashed line.

#### 570 3.2 COS mixing ratios above and within the canopy

571 While the canopy depleted the ambient  $\chi_{COS}$  during day as well as nighttime, we found that the  $\chi_{COS}$  reached values as low as 572 134 ppt (depletion of 102 ppt with respect to the mixing ratio at canopy height) during nighttime (see Fig. 3) at the bottom of 573 the canopy in contrast to the midday  $\chi_{COS}$ , which only went down to 389 ppt (depletion of 125 ppt with respect to the mixing 574 ratio at canopy height) served a decrease in  $\gamma_{CO2}$  (up to 26 ppm) within the most upper layers of the canopy compared 575 to  $\chi_{CO2}$  at canopy height during daytime, while  $\chi_{CO2}$  increased within the lowest layers compared to  $\chi_{CO2}$  at the canopy height 576 due to soil respiration. The above canopy  $\chi_{cos}$  increased considerably starting at the onset of the day and reached 587 ppt at 577 4 p.m. with a steep increase until 11 a.m. Over the course of the season the midday ambient  $\chi_{COS}$  decreased from 500 ±28 ppt 578 from mid-June to mid-July to 405±29 ppt in October with the trend of increasing  $\chi_{COS}$  starting at the end of September (see 579 Fig. S6).



580



#### 586

#### 587 3.2-3\_COS soil flux

The fluxes resulting from the soil chamber measurements ranged from -6.3 to 40.9 pmol m<sup>-2</sup>s<sup>-1</sup>, with positive fluxes denoting emission (see Fig.  $\frac{3-4}{2}$  panel d).

During nighttime (RSW = 0, n = 43), 74.4 % of the COS fluxes were negative, implying that the soil of the grassland acted 590 as a net sink for COS During nighttime (R<sub>SW</sub>=0, n = 43), the soils of the grassland acted as a net sink for COS 74.4 % of the 591 time (range of -4.4 to 6.9 pmol m<sup>-2</sup>s<sup>-1</sup>), whereas soils transitioned to a source in 88.5 % of all daytime measurements ( $R_{SW}$  > 592 0, n = 200), reaching the highest fluxes of 40.9 pmol  $m^2s^{-1}$  during midday (see Fig. 3-4 a-c and Fig. S3). This diel pattern 593 was maintained over the course of the season, however with decreasing maximum COS source strength of the soil towards 594 the end of the season (Fig. 3-4 a-c and Fig. S3). The random forest regression revealed that the most important variable for 595 predicting the soil fluxes was the incident shortwave radiation reaching the soil surface (R<sub>SW-soil</sub>), accounting for more than 596 597 73.53 % of the total variance explained by the final model, while SWC and T<sub>soil</sub> only accounted for 17.84 % and 8.62 %, respectively. The fast response of the COS soil fluxes to changes in  $R_{SW}$  can be seen in Fig. 3-4 a, where we observed a 598 599 decrease of R<sub>SW-soil</sub> as well as the COS soil flux during a cloudy period, even when the soil temperature still increased. Soil fluxes estimated with the random forest regression ranged from -1.3 to 5.0 pmol m<sup>-2</sup>s<sup>-1</sup>, reflecting the fact that under real-600 world conditions very little solar radiation reaches the soil surface. (Fig. 3-4\_e). The resulting emissions peaked during 601 602 daytime shortly after the cuts when a high proportion of incident radiation was reaching the soil surface, while simulated nighttime fluxes were dominated by uptake (in 93 % of all cases) for the whole season. 603



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#### 613 3.34 COS and CO<sub>2</sub> ecosystem-scale fluxes

614 The grassland acted as a net sink for COS during the majority of our study period with 80 % of the COS ecosystem fluxes between  $-\frac{60.2}{56.0}$  pmol m<sup>-2</sup>s<sup>-1</sup> and  $-\frac{12.5}{4.5}$  pmol m<sup>-2</sup>s<sup>-1</sup> during daytime and  $-\frac{41.5}{37.8}$  pmol m<sup>-2</sup>s<sup>-1</sup> and  $-\frac{4.6}{9.2}$  pmol m<sup>-2</sup>s<sup>-1</sup> 615 during nighttime. However, wWe also observed a net release of COS at the field site 4.5-11.2 % of the time. The net CO<sub>2</sub> 616 fluxes ranged from -20.74 to  $\frac{3.24.8}{2.24.8}$  µmol m<sup>-2</sup>s<sup>-1</sup> and  $\frac{1.6}{4.6}$  and  $\frac{1.6}{28.736.4}$  µmol m<sup>-2</sup>s<sup>-1</sup> for 80% of all observation during day 617 618 and nighttime, with daytime net emissions occurring after the cuttings of the grassland (Fig. 2 a-c and Fig. 4-5 a). While the 619 COS nighttime fluxes remained unaffected by the cuts (Fig. 45 c), the daytime fluxes showed a high variability (see Fig. 45620 b). Especially after the cuts we observed a strong decline in COS uptake (Fig. 4 b) and the grassland even turned into a net source for COS in middays (Fig. 2 a-c) with a highest emission flux of 26.8 pmol m<sup>-2</sup>s<sup>-1</sup> (midday median) in August after the 621 622 cut. Especially after the cuts we observed a strong decline in COS uptake and even times where the grassland turned into a to 24.5 pmol m<sup>-2</sup>s<sup>-1</sup> (Fig. 4 b) We observed COS emissions for up to 8 days 623 624 after the cut, when the dried litter had already been removed (Fig. 2 a-c). Compared to respiration processes outpacing GPP 625 almost instantaneously after the cuts, the grassland reached its peak COS emission on the day of the cut only in July, 626 whereas the peak was reached five days after the cut in June and August (Fig. 2 a-c). The cut in October led to a reduction in 627 COS uptake, which declined across several days and did not recover, as the end of the season was reached (Fig. 2 d & Fig. 5 b). The cut in October led to a reduction in COS uptake, which was lowest three days after the cut (Fig. 2 d). After the 628 629 fertilization of the field in October the grassland also turned into a source for COS during midday hours for one day (Fig. 45 630 b). Our flux measurements also included a time when the grassland was covered with snow (on the 20.05.2015), which reduced the COS (and CO<sub>2</sub>) fluxes to values close to zero. Over the course of the season, we observed a decline in the 631 magnitude of the daytime COS uptake from  $-50.69 \pm 2524.0-6$  pmol m<sup>-2</sup>s<sup>-1</sup> during midday in the first week of May down to -632  $\frac{29.610.3}{25.510.4}$  pmol m<sup>-2</sup>s<sup>-1</sup> in the last week of October, which was also correlated with the decline in the CO<sub>2</sub> sink 633

- 634 strength and shift to net emission of  $CO_2$ -from -19.9 ± 8.0 µmol m<sup>-2</sup>s<sup>-1</sup> to  $\frac{11.9 4.4}{2} \pm \frac{36.91.5}{36.91.5}$  µmol m<sup>-2</sup>s<sup>-1</sup> (Fig. 4-5 a-eb). We 635 observed an increase in COS and CO<sub>2</sub> fluxes within the growing phases after the cuts only up to an LAI of ~ 4 (-) (Fig. S4-
- 636 S5), which then levelled out for COS and declined for CO<sub>2</sub> due to ecosystem respiration compensating GPP.



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**Figure 45**: Seasonal cycle of the half hourly  $CO_2$  (a), COS daytime (b) and COS nighttime (c) ecosystem fluxes in  $\mu$ mol m<sup>-2</sup>s<sup>-1</sup> and pmol m<sup>-2</sup>s<sup>-1</sup> depicted by black circles if they are above the limit of detection (LOD) and grey x's if they are below (Langford et al., 2015). The red circles depict the mean fluxes between 11 a.m. and 2 p.m. CET for (a & b) and between 11 p.m. and 2 a.m. for (c) that are above the LOD, while the red x's indicate means below the LOD. The red error bars depict the ±1standard deviation of the mean. The blue lines depict the running mean (5 days) for the mean fluxes. The black dashed lines depict the cuttings of the grassland.

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The seasonal pattern of a decrease in COS sink strength was similar for nighttime fluxes (-18.4<u>0</u>± 29.7<u>-6</u> pmol m<sup>-2</sup>s<sup>-1</sup> to -13.010.6 ± 22.518.2 pmol m<sup>-2</sup>s<sup>-1</sup>) (Fig. 4a<u>5c</u>). The mean nighttime respiration also decreased over the course of the season from 15.9 ± 28.2 pmol <u>umol</u> m<sup>-2</sup>s<sup>-1</sup> to <u>12.9.4</u> ± <u>31.717.5 pmol <u>umol</u> m<sup>-2</sup>s<sup>-1</sup> between May and October (Fig. 5a).</u>

647 Periods between May and August of low (after cuts) and high (before cuts) LAI were compared as diel courses (Fig. 5). Over 648 the course of the day, both periods were characterized by a mean uptake of COS (Fig  $\frac{5-6}{2}$ c & d). Even though the uptake was similar during nighttime, the daytime pattern differed considerably. The modelled contribution of the soil to the ecosystem 649 scale COS flux under high LAI conditions (Fig.  $\frac{5-6}{2}$ ) was minor, contributing between 1.3 % and 5.5/5.7 % of the 650 ecosystem flux during midday and morning/evening, respectively. In contrast, during low LAI conditions the soil 651 652 contribution to the ecosystem fluxes increased during daytime and contributed up to 8082.54% of the mean hourly COS ecosystem flux (Fig 56. c). While the grassland acted as a stronger sink for COS during daytime at a high LAI, reaching peak 653 mean uptake values of up to -41.8 pmol m<sup>-2</sup> s<sup>-1</sup> $\pm$  16.8 pmol m<sup>-2</sup> s<sup>-1</sup> during midday, the mean daytime sink strength weakened 654 and we observed close to zero fluxes during midday in periods of low LAI. The magnitude of the soil flux  $(2 \pm 1 \text{ pmol m}^2 \text{ s}^2)$ 655 <sup>1</sup>) was not high enough to explain the difference variation of up to  $\frac{-23.726.0}{23.726.0}$  pmol m<sup>-2</sup>s<sup>-1</sup> between the measured COS 656 ecosystem flux and COS flux resulting from the FP+ model (Fig 5-6 c), suggesting a missing COS source. For phases of high 657 LAI we saw a good agreement between hourly averaged modelled and measured COS ecosystem fluxes (Fig 5-6 d). While 658 659 the grassland acted as a net sink for  $CO_2$  during periods of high LAI (Fig. 5-6 b), a combination of a decline in GPP and an 660 increase in daytime RECO, as more incoming radiation was heating the soil surface, turned it into a net source during midday in periods of low LAI (Fig. 5-6 a). 661 662





**Figure 56.** Mean diel variation of the measured and modelled CO<sub>2</sub> (a & b) and COS (c & d) fluxes for phases of low (LAI <=3) (a & c) and high (LAI >= 4) (b & d) from May to August. The carats depict the modelled gross primary productivity (blue), the modelled ecosystem respiration (red) and the measured CO<sub>2</sub> ecosystem fluxes (black) in µmol m<sup>-2</sup>s<sup>-1</sup>. The circles depict the modelled COS soil flux (yellow), the modelled COS ecosystem flux (turquoise) and the measured  $CO_2$ -COS ecosystem fluxes (black) in pmol m<sup>-2</sup>s<sup>-1</sup>. The red area depicts the difference between the measured cosystem flux and the sum of the modelled fluxes. The grey areas depict the ±1 standard deviation of the mean for all the measured fluxes. The white bars depict the diel mean total incoming shortwave radiation (W m<sup>-2</sup>s<sup>-1</sup>) while the grey bars indicate the diel mean shortwave radiation reaching the soil surface.

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#### 672 **3.4 COS mixing ratios above and within the canopy**

673	While the canopy depleted the ambient $\chi_{cos}$ -during day as well as nighttime, we found that the $\chi_{cos}$ -reached values as low as
674	134 ppt (depletion of 102 ppt with respect to the mixing ratio at canopy height) during nighttime (see Fig. 6) at the bottom of
675	the canopy in contrast to the midday $\chi_{COS}$ , which only went down to 389 ppt (depletion of 125 ppt with respect to the mixing
676	ratio at canopy height). We observed a decrease in $\chi_{CO2}$ (up to 26 ppm) within the most upper layers of the canopy compared
677	to $\chi_{co2}$ at canopy height during daytime, while $\chi_{co2}$ increased within the lowest layers compared to $\chi_{co2}$ at the canopy height
678	due to soil respiration. The above canopy $\chi_{COS}$ -increased considerably starting at the onset of the day and reached 587 ppt at
679	16:00. Over the course of the season the midday ambient $\chi_{COS}$ -decreased from 500 ±28 ppt from mid-June to mid-July to
680	$405\pm29$ ppt in October with the trend of increasing $\chi_{\cos}$ starting at the end of September (see Fig. S6).



D.



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#### 688 3.5 Leaf and ecosystem relative uptake

689 The LRU at high-light conditions, 1, which we calculated using the FP+ algorithm increased from relatively stable precut levels of 0.9-1.4 (-) after before the  $2^{nd}$  and the  $4^{th}$ - $1^{st}$  cut to up to 1.5-6 (-) after the  $4^{th}$  cut (Fig. 7a). After the decrease in  $\iota$ 690 between the 2<sup>nd</sup> and the 3<sup>rd</sup> cut, 1 increased steadily until the 4<sup>th</sup> cut, with the 3<sup>rd</sup> cut seemingly not having an effect. The 691 reason for the increase in 1 after the 2nd and 4th cut was a stronger decrease in GPP than the COS uptake, while both 692 decreased more evenly after the 3<sup>rd</sup> cut (Fig. 7b). We observed 1 in the period before the 4<sup>th</sup> cut to be influenced not only by a 693 decrease in COS uptake, but also by a decrease in COS mixing ratio (Fig 7b). The mean midday ERUs varied between 694 695  $\frac{1.92.0}{2.0} \pm 0.1(-)$  before and  $4.56 \pm 0.34$  (-) after the cuts-when excluding and 3.9including the first cut. The larger difference between the ERU and Lafter the cuts reflect that we observed similar respiration rates at low and high LAI 696 697 (Fig 6a-b). The larger difference between the ERU and tafter the cuts reflect the higher respiration rates of the ecosystem. 698 Under low light conditions, the LRU increased during pre- and post-cut phases in a similar manner with the last 15-day

period in October showing an earlier increase in the morning and evening (see Fig. S7). 699

<sup>10</sup>[a 6.5ERU 6 LAI 5.5ι un rtaint  $\mathbf{5}$ 5 (-) and ERU (-) 4.54 ₹ 3.53 2.52 0.5 150  $\frac{1.5}{300}$ 200 250Day of the year 2015 b  $F_{COS}$   $\chi COS$  GPP  $\chi CO2$ sum (E δ1

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**Figure 7.** (a) The seasonal cycle of t (black line) with the 95% confidence interval (gray area) resulting from the FP+ model and the midday mean (11 a.m. -2 p.m. at PAR > 800 µmol m.<sub>2</sub> s.<sub>1</sub>) ecosystem relative uptake (ERU) (blue line) using the CO<sub>2</sub> ecosystem flux for the calculation windows (~15 days adjusted to cuts). The dashed black line depicts the progression of the leaf area index (LAI) of the grassland. (b) The contribution of the drivers (F<sub>COS</sub>,  $\chi_{COS}$ , GPP and  $\chi_{CO2}$ ) to the changes in t between all calculation windows and the reference period (DOY 169-188) resulting from the linear perturbation analysis compared to the observed change in t ( $\delta$ 1).

#### 707 4 Discussion

#### 708 4.1 COS mixing ratios

709 The continuous seasonal decrease in above-canopy  $\chi_{COS}$  fr 500 pp tober) was within the 710 range of published records observing mixing ratios to decrease from 465 (in summer) to 375 ppt (in winter) (Kuhn et al., 711 1999). This pattern is typical for the northern hemisphere and the COS drawdown by terrestrial ecosystems (Montzka et al., 712 2007). We found the lowest  $\chi_{COS}$  at the end of September, which coincides with the lowest ambient mixing ratios of COS, 713 measured in Ireland, the closest COS observation site Mace Head (MHD) of NOAA, on the 6<sup>th</sup> of October (Fig. S6). 714 diurnal cycle revealed a continuous decrease of Y co The Gradient atmosphere (> 500ppt) down 715 during nighttime.extremely low COS canopy mixing ratios we observed within the canopy, like 716 this have also been reported by Rastogi et al. (2018), who measured a mean  $\chi_{COS}$  minimum 717 of 152 ppt at 1 m above the soil within an old growth forest. Compared to the consistent decrease of COS below the canopy 718 level during day and nighttime, the gradient for CO<sub>2</sub> reverses during nighttime due to ongoing respiration processes while 719 plants are not photosynthetically active. Even though the COS mixing ratio at the layer closest to the soil were higher during 720 day than during nighttime, the absolute decrease in COS was lower during nighttime due to partial stomatal closure 721 (Kooijmans et al., 2017;Campbell et al., 2017). The absolute difference in concentrations during day and nighttime originate 722 from changes in the height of the planetary boundary layer (PBL). While the PBL is shallow during nighttime and the COS 723 mixing ratio decreases due to sink strength of the grassland, at the onset of the day, the PBL layer height increases fast and 724 COS rich air is transported down to the ecosystem (Fig. S12) (Campbell et al., 2017). A similar steep increase until midday 725 has also been observed by Rastogi et al. (2018). Even though CO<sub>2</sub> and COS share a similar pathway into plants, reflected by 726 their respective decrease in the mixing ratios within the canopy, we saw a difference at the lower levels of our gradient 727 analysis during daytime. We only observed an increase in CO2 mixing ratios, caused by the release of CO2 through 728 respiration processes in the soil, whereas COS mixing ratios further declined down to the soil surface. This supports our soil

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model, which predicted only minor COS fluxes under conditions of high LAI, when only a small portion of incident
 radiation reaches the soil surface.

#### 731 4.1-2 Soil fluxes

The nighttime soil chamber measurements compare well in terms of magnitude with the COS fluxes resulting from studies using dark chambers in agricultural and grassland sites (Whelan et al., 2018;Maseyk et al., 2014;Whelan and Rhew, 2016;Liu et al., 2010) and indicate the soil to be a small sink for COS. The current understanding of COS soil exchange links the COS consumption to soil biota e.g. bacteria and fungi, possessing the ubiquitous enzyme CA (Kesselmeier et al., 1999;Meredith et al., 2019). However, we also found 12 % of all nighttime fluxes to be emission. The origin of COS in soils on the other hand is still highly debated, but comparisons of untreated and sterilized soils suggest yet unknown abiotic processes (Meredith et al., 2019;Kitz et al., 2019).

739 During daytime, the soil inside the chambers emitted COS at rates of up to 40.9 pmol m<sup>-2</sup>s<sup>-1</sup>. These The high COS emissions 740 resulting from the soil chambers during daytime rates-lie at the upper end of recently stated values of agricultural and 741 grassland sites (Whelan et al., 2018;Kitz et al., 2017;Maseyk et al., 2014;Liu et al., 2010). Partly, this can be attributed to the 742 type of chambers we used and their deployment. We allowed the full spectrum of incoming radiation to reach the soils 743 surface, whereas most other studies used dark chambers. Therefore we were able to capture the influence of COS emission 744 processes coupled to thermo- and photo production on our COS soil fluxes (Whelan and Rhew, 2015;Kitz et al., 745 2019;Meredith et al., 2018). This also led to lower peak soil emissions of COS at the end of the season, when the incoming 746 radiation declined.

The low COS mixing ratios observed in the lowermost canopy layers just above the soil surface emphasize the importance of
 using air from within the canopy for soil chamber measurements and not COS richer air from above the canopy, which
 would increase the COS gradient and thus increase the uptake/decrease emission of COS to/from the soil.

Our modelled COS soil fluxes peak at about 12% of the maximum emissions retrieved from the soil chambers. This is owed to the difference in incident radiation reaching the soil surface between the fluxes resulting from chamber measurements and our model. For the chambers, the aboveground biomass was removed, whereas our modelled fluxes were adjusted for undisturbed canopy conditions. In the gradient mixing ratio data, during pre-cut conditions, we also did not see an increase in COS mixing ratio within the canopy, which would have been a hint for a soil COS source.

Another factor contributing to the high COS soil emissions might be the yearly fertilization using slurry, as high nitrogen content in soils has been linked to a higher source strength of COS (Kaisermann et al., 2018). This agrees well with the study of Kitz et al. (2019), who found a correlation between increased soil nitrogen content and soil COS emission in a laboratory experiment with samples taken from the grassland at two different dates (i.e. June and September).

#### 760 4.2-3 Ecosystem fluxes

Our observations show that the agriculturally used grassland acted as a major sink for COS during the growing season. The fluxes fit well within or even exceeded the COS uptake rates of published grassland and agricultural sites during their growing phases (Billesbach et al., 2014;Whelan and Rhew, 2016;Geng and Mu, 2004). The late snow event that occurred in the peak growing season almost completely inhibited the exchange of  $CO_2$  and COS, as the snow acted as a diffusion barrier for these compounds (Björkman et al., 2010).

The cuttings and the consecutive drying of the above ground plant material at the site had a major influence on the COS exchange. During these events the grassland turned into a source for  $CO_2$  and COS. COS emissions of a similar magnitude

768 This hasve also been reported at agricultural fields in phases of senescence (Maseyk et al., 2014;Billesbach et al., 2014).

769 Although the soil was a strong source for COS, caused by the high  $R_{soil}$  and  $T_{soil}$  (Whelan and Rhew, 2015;Kitz et al.,

770 2019;Meredith et al., 2018), and the sink strength of the grassland was low due to the reduced aboveground biomass, soil 771 fluxes did not explain the emission on ecosystem level (see Fig. 546a). As plants contain precursors involved in COS 772 emission processes, e.g. methionine and cysteine (Meredith et al., 2018), the plant litter and dying plant parts remaining at 773 the site after the cuts might be the missing source of COS. Laboratory tests of the soil of the grassland have shown that a mixing of dried litter and soil lead to a strong but short-lived emission peak of COS (Kitz et al., 2019). We did not observe 774 775 strong COS emissions after the last cut, as the incoming solar radiation, which we hypothesize to amplify the degradation of 776 sulfur containing compounds of plants, was reduced at the end of the season. Alternatively, the cutting of the grassland might 777 induce stress mediated COS production in the remaining living plant parts (Bloem et al., 2012;Gimeno et al., 2017). The 778 delay in the peak COS emissions at ecosystem scale after the cuts could indicate that some yet unknown biotic or abiotic processes take several days to release COS. 779

#### 780 We also observed another COSThe short-lived timed COS emission by yet unknown biotic or abiotic processes event shortly

after the fertilization of the grassland towards the end of the growing season-<u>was likely triggered by T</u>the increase of

- available nitrogen (Kaisermann et al., 2018) and COS precursors introduced to the soil in the form of cattle slurry (Hörtnagl
   et al., 2018), might have triggered the COS emission by biotic or abiotic processes.
- Due to the independence of CA to catalyze COS without  $R_{PAR}$  (Stimler et al., 2011), the grassland remained a sink for COS during nighttime. Again, the soil sink was too small to explain the total COS exchange (Fig. 56), which indicates that the plant stomata were not fully closed (Kooijmans et al., 2017) and were responsible for the majority of the COS uptake. The minimum or residual stomatal conductances at the field site in Neustift have been reported to be between 10 and 65 mmol m<sup>-</sup> 788  $^{2}$  s<sup>-1</sup> depending on the species (Wohlfahrt, 2004).
- The large variability in COS nighttime fluxes (Fig. 5c) is due to the combination of low wind speeds and stable stratification,
   which results in highly intermittent CO<sub>2</sub> (Wohlfahrt et al., 2005) and COS fluxes compared to daytime. On half-hourly basis,
   even a nighttime net uptake of CO<sub>2</sub> has been reported at the field site, which is typically compensated for by large CO<sub>2</sub>
   emissions in a subsequent averaging period (Wohlfahrt et al., 2005). We also observed this pattern for COS.
- Although we observed phases of high VPD and low SWC (Fig. 1), they did not lead to a decrease in  $CO_2$  and  $CO_3$  ecosystem fluxes (Fig. S1-S2), which has already been observed for the grasslands  $CO_2$  and  $H_2O$  fluxes between 2001 and 2009. The species located at the site were insensitive to progressive drought conditions (Brilli et al., 2011).

#### 796 4.3 COS mixing ratios

797	The continuous decrease in above canopy $\chi_{cos}$ from -500 ppt (in May) to -400 ppt (in October) is within the range of
798	published records observing mixing ratios to decrease from 465 (in summer) to 375 ppt (in winter) (Kuhn et al., 1999). This
799	pattern is typical for the northern hemisphere and the COS drawdown by terrestrial ecosystems (Montzka et al., 2007). We
800	found the lowest X <sub>COS</sub> at the end of September, which coincides with the lowest ambient mixing ratios of COS, measured in
801	Ireland, the closest COS observation site Mace Head (MHD) of NOAA, on the 6 <sup>th</sup> of October.
802	Gradient observations of the diurnal cycle revealed a continuous decrease of $\chi_{COS}$ from the atmosphere (> 500ppt) down to
803	the soil reaching very low concentrations mixing ratios of 134 ppt during nighttime. Low values like this have also been
804	reported by Rastogi et al. (2018), who measured a mean $\chi_{COS}$ -minimum of 152 ppt at 1 m above the soil within an old growth
805	forest. The difference in concentrations during day and nighttime originates from changes in the height of the planetary
806	boundary layer (PBL). While the PBL is shallow during nighttime and the COS mixing ratio decreases due to sink strength
807	of the grassland, at the onset of the day, the PBL layer height increases fast and COS rich air is transported down to the
808	ecosystem. Even though CO2 and COS share a similar pathway into plants, reflected by their respective decrease in the
809	mixing ratios within the canopy, we saw a difference at the lower levels of our gradient analysis during daytime. We only
810	observed an increase in CO <sub>2</sub> -mixing ratios, caused by the release of CO <sub>2</sub> -through respiration processes in the soil, whereas

- 811 COS mixing ratios further declined down to the soil surface. This supports our soil model, which predicted only minor COS
- 812 fluxes under conditions of high LAI, when only a small portion of incident radiation was hitting the soil surface.

#### 813 4.4 LRU

#### 814 The parameter t varied between 0.9 (0.8 1.0) ( ) and 1.5 (1.2 1.8) ( ) during the campaign, where cuts of the grassland tended

815 to result in higher values and of places this study is placed at the lower end of a recent compilation of all-published leaf-level

816 LRUs, that put 95% of all data between 0.7 (-) and 6.2 (-) with a median of 1.7 (-) (Whelan et al., 2018) and also lower than

the LRU of 2.53 (-) estimated for grasslands by Seibt et al. (2010). Even the higher 1 after the cuts was low compared to

818 these studies. The seasonal trend of the LRUs 1 was strongly influenced by the cutting of the grass and can be attributed

819 mainly to changes in the ratio of COS uptake to GPP. However, we also observed a strong decline in the ambient mixing

ratio of COS, which also had an equally strong influence on the change in t as the COS flux for the 15 day window beforethe last cut (Fig 7 b).

822 Even though the changes in t can be explained, it is important to keep in mind that the grassland was a source for COS on

823 ecosystem level after the cuts. For the calculation of LRUs we had to remove those the canopy flux data containing COS

824 and/or CO<sub>2</sub> emissions observations from the data-since they would yield negative values for ERU and LRU (see Eq.8). This

825 indicates that the unknown source strength after cuts likely decreases the post-cut 1's.

#### 826 5 Conclusion

827 Due to the management interventions at the grassland site, the leaf area development was decoupled from seasonal changes 828 in environmental forcing. This allowed us to measure concurrent CO2 and COS fluxes at soil and ecosystem level for 829 multiple growing periods within one season. The LAI on seasonal scale as well as incoming solar radiation on hourly to 830 seasonal scales determined whether soils were a source or a sink for COS. The incoming shortwave radiation reaching the 831 soil surface had a decisive influence on the COS soil surface flux and thus supports our hypothesis H4. The covariance 832 between the daytime CO<sub>2</sub> and COS fluxes on daily to seasonal level was high and the fluxes only diverged after the cuts, 833 leading to higher LRUs. Beside the perturbations of the ecosystem, the sink strength of the grassland was high for COS and 834 declined over the course of the season (H1). The COS emissions at ecosystem scale shortly after the cuts, which could not be 835 explained by the soil source, raise questions about other unknown mechanisms of COS production within ecosystems (H2). 836 With the exception of short periods after the cuts, the LRUs under high light conditions were relatively constant during the season, indicating a good correlation between the COS flux and GPP under stable conditions (H3). 837

#### 838 6. Data availability

Bata and materials availability: <u>https://doi.org/10.5281/zenodo.3886554</u>
 Will be uploaded to https://zenodo.org/.
 840

#### 841 7. Author contributions

Felix M. Spielmann: Data curation, Formal analysis, Investigation, Methodology, Software, Visualization, Writing – original
 draft

844 Albin Hammerle: Data curation, Investigation, Software, Writing - original draft

- 845 Florian Kitz: Data curation, Formal analysis, Investigation, Methodology, Software, Writing original draft
- 846 Katharina Gerdel: Investigation, Software, Writing original draft

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847 Georg Wohlfahrt: Conceptualization, Funding acquisition, Investigation, Methodology, Project administration, Software,

848 Supervision, Writing - original draft

#### 849 8. Competing interests

850 The authors declare no competing financial interests.

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858 interests.

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