

1 Dynamics of short-term ecosystem carbon fluxes induced by 2 precipitation events in a semiarid grassland.

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15 **Abstract.** Infrequent and small PPT events characterize precipitation (PPT) patterns in semiarid grasslands;
16 however, plants and soil microorganisms are adapted to use the unpredictable small pulses of water. Several
17 studies have shown short-term responses of carbon and nitrogen mineralization rates (called the priming effect
18 or the Birch effect) stimulated by wet-dry cycles; however, dynamics, drivers, and the contribution of the
19 “priming effect” to the annual C balance are poorly understood. Thus, we analysed six years of continuous net
20 ecosystem exchange measurements to evaluate the effect of the PPT periodicity and magnitude of individual
21 PPT events on the daily/annual ecosystem C balance (NEE) in a semiarid grassland. We included the period
22 between PPT events, previous daytime NEE rate, and previous soil moisture content as the main drivers of the
23 priming effect. Ecosystem respiration (ER) responded within few hours following a PPT event, whereas it took
24 five-nine days for gross ecosystem exchange (GEE; where $-NEE = GEE + ER$) to respond. Precipitation events
25 as low as 0.25 mm increased ER, but cumulative PPT > 40 mm infiltrating deep into the soil profile stimulated
26 GEE. Overall, ER fluxes following PPT events were related to the change of soil water content at shallow depth
27 and previous soil conditions (e.g., previous NEE rate, previous soil water content) and the size of the stimulus
28 (e.g., PPT event size). Carbon effluxes from the priming effect accounted for less than 5% of ecosystem
29 respiration but were significantly high respect to the carbon balance. In the long-term, changes in PPT regimes
30 to more intense and less frequent PPT events, as expected by the effects of climate change, could convert the
31 semiarid grassland from a small C sink to a C source.

32 Keywords: Eddy covariance, net ecosystem exchange, ecosystem respiration, *Bouteloua gracilis*, blue grama,
33 priming effect, Birch effect.

35 1. Introduction

36 Arid lands comprise many ecosystem types covering more than 30% of terrestrial land (Lal, 2004). In these
37 ecosystems annual potential evapotranspiration is larger than yearly precipitation due to regional atmospheric
38 high-pressure zones (i.e., Hadley cells), continental winds, cold oceanic winds and local orographic effects that
39 reduce the precipitation amounts (Maliva and Missimer, 2012). Here, precipitation (PPT) occurs as infrequent,
40 discrete, small (< 5 mm), and unpredictable events (Noy-Meir, 1973; Loik et al, 2004). This results in water-
41 limited ecosystems, where biological activity is restricted to periods of soil water availability (Lauenroth and
42 Sala, 1992). Consequently, the productivity and stability of these ecosystems are more vulnerable to changes in
43 climate, particularly to changes in the historic mean annual PPT amounts (MAP; Wang et al., 2021) and the
44 change in the periodicity (i.e., frequency) of these PPT events (Korell et al., 2021; Nielsen and Ball, 2015).

45 Precipitation stimulates short-term changes of carbon and nitrogen mineralization rates because soil
46 microorganisms activate with increased soil water content (Turner and Haygarth, 2001). This “priming effect”
47 (Borken and Matzner, 2009), also called the Birch effect (Birch, 1964), describes the soil carbon released from
48 the decomposition of heterotrophic sources to the atmosphere following soil rewetting. The amount and timing
49 of PPT events modify the magnitude and duration of the priming effect by modulating soil wet-dry cycles. The
50 size of a PPT event determines the temporal duration and the biotic components that respond to the pulse
51 (Huxman et al., 2004a), thus defining the magnitude and direction of CO₂ effluxes (Chen et al., 2009). In
52 general, small precipitation events that induce changes in soil humidity at shallow depths do not induce plant
53 activity, but activate soil microorganisms (Collins et al., 2008) and consequently enhance CO₂ effluxes (Vargas
54 et al., 2012). On the other hand, successive rewetting cycles reduce carbon mineralization rates as the amount
55 of available organic labile carbon declines (Jarvis et al., 2007). Thus, PPT events after long drought periods
56 (until nine months in semiarid grassland) trigger larger and longer soil respiration efflux rates than consecutive
57 PPT events (Reichmann et al., 2013; Vargas et al., 2018).

58 At the ecosystem scale, deserts and grasslands have shown larger CO₂ efflux rates after rewetting than temperate
59 ecosystems or croplands (Kim et al., 2012) and in ecosystems with low soil organic carbon content (Bastida et
60 al., 2019). Characteristics and dynamics of these short-term soil C effluxes were addressed by the “Threshold-
61 Delay” model (T-D model, Ogle and Reynolds 2004). The T-D model takes previous environmental conditions,
62 PPT event size, PPT thresholds, and time-delays to inform the time constants that modulate ecosystem
63 responses after a PPT event. Moreover, Huxman et al. (2004a) described the dynamics of the net ecosystem
64 exchange of carbon (NEE) and its components (gross ecosystem exchange = GEE, and ecosystem respiration
65 = ER, such as $-NEE = GEE + ER$) with parameters of the T-D model (Fig. A1). GEE and ER have different
66 time delays based on threshold PPT quantities and event size, with ER responding to smaller PPT events than
67 GEE (Huxman et al. 2004a). In addition, GEE and ER have asymptotic responses to large PPT events (the upper
68 PPT thresholds), with an upper ER threshold lower than that found for GEE (Huxman et al. 2004).

69 In the semiarid grasslands of Mexico, small PPT events likely activate biological soil crusts (BSC) that cover
70 up to 60% of plant interspaces (Concostrina-Zubiri et al., 2014) and stimulate ER instead of C uptake. However,
71 *Bouteloua gracilis* H.B.K. Lag ex Steud (blue grama), the keystone species in the semiarid grassland of Mexico

72 (Medina-Roldán et al., 2007) may contribute to C uptake because of its adaptations to take advantage of small
73 PPT events (Sala and Lauenroth, 1982, Medina-Roldán et al., 2013). Understanding disturbances of ecosystem
74 processes (C fluxes) due to changing regional PPT patterns in semiarid grasslands are particularly salient given
75 that the global circulation models forecast between 10% and 30% reduction of summer and winter precipitation,
76 respectively, by the end of the 21st Century (Christensen et al., 2007). Furthermore, PPT patterns are expected
77 to have fewer events with more water quantity per event (Easterling et al., 2000).

78 The objective of this study was to evaluate the effect of PPT periodicity and magnitude of individual PPT events
79 and a priori soil moisture conditions on daily and annual ecosystem C balance (NEE) for the semiarid grassland
80 in Mexico. Over a six-year study period, we examined event-based PPT amount, the period between PPT
81 events, and the previous daytime NEE rate and soil water content at two depths as the main drivers of daily
82 mean NEE change rate. Because we were interested in short-term NEE changes and their components, only
83 short-term NEE changes within a few days following a PPT event were evaluated. Effects on daily mean GEE
84 ($GEE = -NEE + ER$) were also evaluated at the beginning of the growing season. Based on the T-D model
85 (Ogle and Reynolds, 2004), we expect that; 1) semiarid grassland will exhibit a quick response (short time-
86 delay) to small PPT events (Low PPT threshold) through positive NEE fluxes (C release, H1). Moreover, 2)
87 ER and GEE (C release and C uptake, respectively) will differ in their response times and PPT thresholds, with
88 shorter time-delays and lower PPT thresholds for ER than GEE (H2). This response is because small PPT
89 events should enhance ER mainly through heterotrophic respiration of soil surface microorganisms that are
90 activated within one hour after wetting (Placella et al., 2012), whereas larger PPT events are required to reach
91 roots at deeper soil profiles and longer times for plants to start growing. On the other hand, we expect that 3)
92 the size and timing of PPT patterns will modulate the magnitude of C efflux; therefore, large precipitation events
93 after long dry periods will release more CO₂ than small or consecutive PPT events (H3). Finally, we expect 4)
94 C efflux after PPT events will be a meaningful CO₂ source to the atmosphere in the semiarid grassland,
95 decreasing the ecosystem's annual net C uptake (H4).

96 **2. Materials and methods**

97 **2.1 Site description**

98 The study site is located on a shortgrass steppe within the Llanos de Ojuelos subprovince of Jalisco state,
99 Mexico. The shortgrass biome in Mexico extends from the North American Midwest along a strip that follows
100 the Sierra Madre Occidental through the Chihuahuan Desert into the sub-province Llanos de Ojuelos.
101 Vegetation is dominated by grasses, with *Bouteloua gracilis* (Willd. ex Kunth) Lag. ex Griffiths as the key
102 grass species, forming near mono-specific stands. The region has a semiarid climate with mean annual
103 precipitation of 424 mm ± 11 mm (last 30 years, Delgado-Balbuena et al., 2019) distributed mainly between
104 June and September and with 6 – 9 moths of low PPT. Winter-summer rain accounts for < 20% of annual
105 precipitation (Delgado-Balbuena et al., 2019). The mean annual temperature is 17.5 ± 0.5 °C. The topography
106 is characterized by valleys and gentle rolling hills with soils classified as haplic xerosols (associated with
107 lithosols and eutric planosols), and haplic phaeozems (associated with lithosols) (Aguado-Santacruz, 1993).

108 Soils are shallow, with average depth of 0.3-0.4 m containing a cemented layer at ~ 0.5 m deep, with textures
109 dominated by silty clay and sandy loam soils (Aguado-Santacruz, 1993).
110 The study site is a fenced area of ~64 ha of semiarid grassland under grazing management. A 6 m high tower
111 was placed at the center of the area of interest to support carbon-energy flux measurements and meteorological
112 instruments. That location allowed an ever-changing and integrated measurement footprint of 320 m, 410 m,
113 580 m, and 260 m from the tower according to the N, E, S, and W orientations, respectively. The study site is
114 part of the MexFlux network (Vargas et al., 2013).

115 **2.2 Meteorological and soil measurements**

116 Meteorological data were collected continuously at a rate of 1 s and averaged at 30 min intervals using a
117 datalogger (CR3000, Campbell Scientific Inc., Logan, Utah). Variables measured included air temperature and
118 relative humidity (HMP45C, Vaisala, Helsinki, Finland) housed into a radiation shield (R.M. Young Company
119 Inc., Traverse City, MI), incident and reflected shortwave and longwave solar radiation (NR01, Hukseflux,
120 Netherlands), and photosynthetic photon flux density (PPFD, PAR lite, Kipp and Zonen, Delft, the
121 Netherlands). Soil variables were measured at a 5 min frequency and averaged at 30 min intervals. These
122 included volumetric soil water content (CS616, Campbell Sci., Logan, UT) positioned horizontally to 2.5 cm
123 and 15 cm deep, average soil temperature of the top 8 cm soil profile, and soil temperature at 5 cm deep (T108
124 temperature probes, Campbell Scientific Inc., Logan, UT). Soil temperature variables were acquired with
125 another datalogger (CR510, Campbell Scientific Inc., Logan, UT). Precipitation was measured with a bucket
126 rain gauge installed 5 m away from the tower (FTS, Victoria, British Columbia, Canada) at 1 m.a.g.l.

127 **2.3 Net ecosystem CO₂ exchange measurements**

128 An open path eddy covariance system was placed at 3 m high to cover a fetch of 300 m and used to measure
129 NEE over the semiarid grassland. The system consisted of a three-dimensional sonic anemometer (CSAT-3D,
130 Campbell Sci., Logan, UT) for measuring wind velocity on each polar coordinate (u , v , w) and sonic temperature
131 (θ_s) and an open-path infrared gas analyzer (IRGA, Li-7500, LI-COR Inc., Lincoln, NE) to measure CO₂ and
132 water vapor concentrations. Instruments were mounted in a tower at 3 m above the soil surface, oriented
133 towards the prevailing winds. The IRGA sensor was mounted next to—and 10 cm offset from the anemometer
134 transducers, the center of the IRGA optical path was centered with the distance between the vertically oriented
135 sonic transducers and tilted 45° to avoid dust and water accumulation in the IRGA optical path. Digital signal
136 of both sensors was recorded at a sampling rate of 10 Hz in a datalogger (CR3000, Campbell Scientific Inc.,
137 Logan, UT) (Ocheltree and Loescher 2007). NEE was estimated as:

$$138 \quad NEE = \overline{w'CO_2'} \quad (1)$$

139 overbar denotes time averaging (30 min), and primes are the deviations of instantaneous values (at 10 Hz) of
140 vertical windspeed (w' , ms⁻¹) and molar volume of CO₂ (CO_2' , μmol CO₂ m⁻³), from the block-averaged mean.
141 Micrometeorological convention was used, where negative NEE values stand for ecosystem C uptake (Loescher

142 et al., 2006). We did not estimate a storage flux because of the low vegetation stature and well-mixed
143 conditions; therefore, we assumed it would be 0 over a 24-h period (Loescher et al., 2006).

144 **2.4 Data processing**

145 Raw eddy covariance data were processed in EdiRe (v1.5.0.10, University of Edinburgh, Edinburgh UK). Wind
146 velocities, sonic temperature, [CO₂], and [H₂O] signals were despiked (Vickers and Mahrt, 1997), any value
147 larger than six standard deviations into a moving window (5 min) was considered a spike, whereas those values
148 with a deviation larger than eight standard deviations were flagged as outliers. A 2-D coordinate rotation was
149 applied to sonic anemometer wind velocities to obtain turbulence statistics perpendicular to the local streamline.
150 Lags between horizontal wind velocity and scalars were removed with a cross-correlation procedure to
151 maximize the covariance among signals. Carbon and water vapor fluxes were estimated as molar fluxes (mol
152 m⁻² s⁻¹) at 30 min block averages, and then they were corrected for air density fluctuations (WPL correction,
153 Webb et al. 1980). Frequency response correction was done after Massman (2000). Sensible heat flux was
154 estimated from the covariance between fluctuations of horizontal wind velocity (*w'*) and sonic temperature (*θ'_s*).
155 This buoyancy flux was corrected for humidity effects (Schotanus et al. 1983, Foken et al., 2012).
156 Fluxes were submitted to quality control procedures, i) stationarity (<50%), ii) integral turbulence
157 characteristics (<50%), iii) flags of IRGA and sonic anemometer (AGC value<75, Max CSAT diagnostic flag
158 = 63) which are frequently caused by raindrops on the anemometer transducers and IRGA path, iv) screening
159 of flux values into expected magnitudes (±20 μmol CO₂ m⁻² s⁻¹), and v) the *u** threshold was used to filter
160 nighttime NEE under poorly developed turbulence. This threshold was defined through the 99% threshold
161 criterion after Reichstein et al. (2005); it varied seasonally among years around 0.1 m s⁻¹.

162 Temporally integrated estimates are noted throughout this paper. Because GEE cannot be measured directly, it
163 was estimated from light-response curves (see below), whereas ER was determined from i) light-response
164 curves and ii) nighttime NEE data (under PPFD < 10 μmol m⁻² s⁻¹ light conditions). Henceforth, ecosystem
165 respiration derived from light-response curves is denoted as “ER”, and as “nighttime NEE” when derived from
166 nighttime net ecosystem exchange data.

167 For identifying changes induced by PPT events on GEE and ER, daytime and nighttime NEE data on a one day-
168 window was adjusted with a rectangular hyperbolic response function to photosynthetic photon flux density
169 (PPFD; Ruimy et al. 1995).

$$170 \quad NEE = \frac{\alpha * PPFD * \beta}{\alpha * PPFD + \beta} + ER \quad (2)$$

171 where, *α* is the apparent quantum yield (μmol CO₂ m⁻² s⁻¹/ μmol CO₂ m⁻² s⁻¹), *β* is maximum photosynthetic
172 capacity (μmol CO₂ m⁻² s⁻¹), and *ER* is the ecosystem respiration (μmol CO₂ m⁻² s⁻¹). Due to *A_{max}* being
173 calculated to unrealistic “infinite” *PPFD*, we calculated a more realistic maximum photosynthetic capacity
174 (*A₂₅₀₀*), which is maximum photosynthesis at 2500 μmol m⁻² s⁻¹. Changes and transitions from ER-dominated
175 NEE fluxes to C-gain processes (GEE) were verified with the shape of the light response curve.

176 We choose this method instead of standard partitioning procedures (i.e. Reichstein et al., 2005 or Lasslop et al.,
177 2010) because we were interested in detecting changes at one day scale. Both algorithms use data windows

178 larger than one day to estimate some parameters and tend to smooth fast changes in soil respiration like the
179 observed in this study. For visually checking for changes in GEE and ER at diel time step, half-hours of NEE
180 were partitioned by Eq. 2 and then averaged by day.

181 **2.5 Gap-filling procedures and characterization of PPT events**

182 Data gaps shorter than two hours were linearly interpolated, whereas gaps larger than two hours were left as
183 empty data. Only daytime-NEE data were used for most of the analysis because nighttime NEE is subjected to
184 quality problems like poorly developed turbulence. Moreover, if mean NEE is estimated from only a few 30-
185 minute nighttime NEE half-hours, the estimate may be biased if the full night cycle is not represented similarly
186 across days. The NEE-related PPT events were selected for analysis based on data quality and availability to
187 evenly cover the daytime cycle (on average more than 85% of NEE data) and then averaged through the day.
188 The daytime-scale was selected to avoid confounding diurnal NEE variability and to achieve robust analyses.
189 All precipitation events between 2011 and 2016 were isolated and filtered by the number of half-hours
190 accounted for mean daily fluxes.

191 Mean ER derived from nighttime NEE data were used for analysis only when more than 50% of the data was
192 available after QA/QC procedures. This data was exclusively used for correlation with environmental and soil
193 data (see statistical analysis section). In contrast, daytime NEE (without partitioning) was used for the analysis
194 of changes in NEE fluxes induced by PPT events.

195 The C flux one day before the PPT event was taken as the reference C flux. The event-response effect (“priming
196 NEE effect”) was measured as the difference between mean daytime NEE post-event and mean daytime NEE
197 pre-event, described as:

$$198 \Delta\text{NEE} = \text{NEE}_{\text{post-event}} - \text{NEE}_{\text{pre-event}} \quad (3)$$

200
201 where NEE is the daytime NEE average ($\mu\text{mol m}^{-2} \text{s}^{-1}$). The same method was used to calculate changes of soil
202 water content at 2.5 and 15 cm depth ($\Delta\text{VWC}_{2.5}$ and ΔVWC_{15} , respectively) and change of photosynthetic
203 photon flux density (ΔPPFD). Intervals between PPT events (hereafter inter-event periods, IEP) were counted
204 in days from the last PPT event, regardless of its magnitude.

205 Enhanced vegetation index (EVI) of 250 m spatial resolution and 8-day time-resolution from NASA’s MODIS
206 instruments (Didan, 2021) was used to approximate plant leaf activity. The Savitzky-Golay (Yang et al., 2014)
207 filter was used to eliminate outliers of EVI derived from adverse atmospheric conditions.

208 Considering that previous conditions are determinant for carbon fluxes, data were divided into “fluxes
209 dominated by photosynthesis (carbon uptake)” and “fluxes dominated by ecosystem respiration (carbon
210 efflux)”. A threshold of $-1 \mu\text{mol m}^{-2} \text{s}^{-1}$ of average previous daytime CO_2 flux was used to divide data. This was
211 done to avoid confounding factors because the environmental drivers of photosynthesis and respiration may
212 differ in magnitude and direction. Moreover, under photosynthetic conditions is hard to identify if a positive
213 change of NEE (less photosynthesis) was due to an increase of soil respiration or a dampening of photosynthesis

214 by less available radiation under cloudy conditions.
215 To estimate the contribution of the priming effect to the annual carbon balance in the semiarid grassland, we
216 averaged and extrapolated Δ NEE by the number of precipitation events per year. Decaying rates, PPT event
217 size, and previous soil and flux conditions were not considered in this approach. Although this is a rough
218 estimation, it provides a broad overview of how precipitation patterns influence the annual carbon balance. It
219 is important to have this broad overview to better understand the impacts of climate change on carbon cycling
220 in semiarid grasslands.

221 **2.6 Statistical analysis**

222 Boosted regression trees analysis (BRT; Elith and Leathwick, 2017) were developed to identify the most
223 important variable controlling this response's priming C effect and thresholds. BRT analysis also was used to
224 identify the form of function, i.e., whether the relationship between independent variables and the priming effect
225 was linear, exponential, sigmoidal, peak from, etc. Independent variables included PPT event size, inter event-
226 periods (IEP), previous, current, and change of volumetric water content (VWC) at two depths (2.5 and 15 cm),
227 soil temperature, previous daytime NEE, enhanced vegetation index (EVI) and change in photosynthetic photon
228 flux density (Δ PPFD). For BRT analysis, data was divided into “photosynthesis dominated” and “respiration
229 dominated” data. On the other hand, to identify delays between C fluxes (ecosystem respiration and gross
230 primary productivity) and precipitation events, a cross-correlation analysis was done. For cross correlation, the
231 parameter of the light response curve was used; the ER was used to identify delays between ecosystem
232 respiration and soil water content at 2.5 cm, and A_{2500} was used to identify delays between gross ecosystem
233 productivity and soil water content at 15 cm, because of ER and A_{2500} were better correlated with soil volumetric
234 water content at 2.5 and 15 cm, respectively. All these variables were detrended before cross-correlation
235 analysis. Finally, linear correlation analyses were performed among environmental variables, priming effect
236 and nighttime NEE (ER), and among independent variables to test for autocorrelations. The “gbm” package
237 (The R core team) was used for performing BRT analysis, whereas the “astsa” package for R was used to
238 conduct cross-correlation analyses.

239 **3. Results**

240 **3.1 Precipitation pattern**

241 Cumulative precipitation for 2011 (288.5 mm) was below the 30-y average for the site (420 mm) and was the
242 worst drought of the last 70-y. In contrast, 2012 received less PPT (393.2 mm), and 2014 and 2016 received
243 more PPT (528.5 and 436 mm, respectively) than average, whereas 2013 (601.6 mm) and 2015 (785.9 mm)
244 were very humid years (Fig. 1). The 6-y differed in precipitation frequency, but they were similar in the size of
245 PPT events with ~60% of the PPT events < 5 mm (Fig. 2a). However, notwithstanding the lower proportion of
246 larger size PPT events (PPT events > 5 mm), they summed similar or even more amount of water than small
247 PPT events (Fig. 2b). Overall, precipitation pattern was characterized by short inter-event periods with 60% of
248 PPT events falling consecutively (IEP < 5 days; Fig. 2c).

249 Soil saturated after large or recurrent PPT events. Largely, soil moisture was maintained at over a 10% in the
250 wettest years, with the largest peak reaching 40% in the summer 2014 (Fig. 1b). Most VWC variability was
251 observed at 2.5 cm rather than 15 cm depth, and it was better correlated with precipitation amount per event (p
252 < 0.05 , $R^2 = 0.72$, Fig. 2d), increasing 0.3 % of VWC per mm of precipitation. The PPT events of 0.25 mm
253 increased the $VWC_{2.5}$ in ~ 1 -2%, but this increase lasted for less than one hour, whereas VWC_{15} increased after
254 PPT ~ 5 mm (data not shown). Additionally, PPT events and soil moisture dynamics at 15 cm depth were out of
255 phase (up to five days between the PPT event and the SWC_{15} peak, Fig. 2e).

256 A total of 391 PPT events were isolated over the six years, but 34% did not accomplish with conditions of diel
257 time representativity ($>85\%$ of NEE data); thus, 256 events were used for statistical analysis. A sample of 100
258 PPT events was used for the respiration dominated fluxes ($>-1.0 \mu\text{mol m}^{-2} \text{s}^{-1}$) and 156 PPT events for the
259 photosynthesis dominated fluxes ($<-1.0 \mu\text{mol m}^{-2} \text{s}^{-1}$). Small precipitation events dominated our dataset but
260 represented well the precipitation pattern of the site. The sample was integrated by events ranging from 0.25
261 to 57.1 mm, and a mean of 5.7 ± 0.53 mm (mean ± 1 SE). Large PPT events occurred after short inter-event
262 periods, and small PPT events were preceded by long inter-event periods. Medium PPT events after long inter-
263 event periods were rare, and extreme large PPT events after long inter-event periods were not observed (Fig.
264 2f).

265 The size of the precipitation event (PPT) and previous soil water content at 2.5 cm depth ($preVWC_{2.5}$) explained
266 a large variation of change in soil water content at 2.5 cm depth ($\Delta VWC_{2.5}$; $R^2 = 0.54$; Fig. 2d). Best correlation
267 among variables was observed between previous soil water content and soil water content at different depths;
268 for instance, VWC_{15} and $pre VWC_{15}$ ($R^2 = 0.84$), between the same variables but at 2.5 cm ($R^2 = 0.81$). The
269 change in NEE (priming effect) did not have a strong relationship with any single variable (Fig. A2).

270 3.2 Time delays and thresholds

271 The minimum PPT event that altered NEE rates was 0.25 mm. Overall, the analysis of half-hour fluxes showed
272 almost instantaneous positive response of NEE to the PPT event that exponentially decreased over time into a
273 half to two hours after the PPT event (Fig. A3). The ER rates increased after 0.25 mm PPT events, but we
274 detected a different threshold for GEE where either a larger PPT event or multiple consecutive events (*e.g.*, $>$
275 40 mm, Fig. 2a) were needed and showed a delay of ~ 5 days after the positive change in VWC at the 15 cm
276 depth, this at the beginning of the growing season (Fig. 3a, b).

277 Cross-correlation analysis of light-response curve parameters showed no lags between ecosystem respiration
278 (ER) and volumetric soil water content at 2.5 cm. (Fig. 3a), whereas there was a lag of 9 days between
279 photosynthetic capacity at 2500 PPFD (A_{2500} ; Fig. 3b) and soil water content, which was larger than the
280 observed at several precipitation events of 2013 (Fig. 2a, b).

281 The BRT analysis showed sigmoidal relationships between the priming effect and environmental variables with
282 different thresholds. At the respiration-dominated period, a minimum change of soil volumetric water content
283 at 2.5 cm affected positively the carbon flux, but a change larger than 8% in this variable did not induce a larger
284 C efflux (upper threshold; Fig. 4). On the other hand, C priming effect was larger under neutral previous NEE

285 (preNEE~0) and decreased in magnitude as preNEE becomes more positive (Fig. 5). Moreover, previous dry
286 conditions at shallow soil depth promoted larger C efflux by the priming effect, and this effect decreased as soil
287 previous conditions were wetter, with a threshold at 15% (Fig. 5). Like the change in soil water content at 2.5
288 cm, even the lowest PPT event (0.25 mm) caused an increase of C efflux, but with a threshold between 10 - 15
289 mm. Precipitation events larger than 15 mm did not enhance the priming effect (Fig. 5). In contrast, in the
290 photosynthesis dominated period, larger priming effect was observed at more negative preNEE ($-7 \mu\text{mol m}^{-2} \text{s}^{-1}$)
291 and had no more effect at $\sim -4 \mu\text{mol m}^{-2} \text{s}^{-1}$. Dry soil conditions enhanced the priming effect at 15 cm depth
292 (< 30%) with a rapid suppression after that. On the other hand, the priming effect was gradually decreasing with
293 reductions in PPFD.

294 Nighttime NEE (ecosystem respiration derived from nighttime NEE data) showed correlation with soil water
295 content at the two depths and EVI; however, the relationship was linear at low soil water content, reached a
296 maximum at medium values of VWC, and then decreased with minimum values at high soil water content. The
297 largest ecosystem respiration was observed at the highest EVI values (Fig. A4).

298 3.3 Dynamics and drivers of the “Priming effect”

299 The priming effect lasted longer with initial larger changes of NEE, i.e., whereas higher was the priming effect
300 (ΔNEE), the C fluxes lasted more time in returning to initial values (before the PPT event); however, decreasing
301 NEE rates were better explained by PPT event size than the initial change of NEE (insert Fig. 4). For instance,
302 after a 13.7 mm PPT event and initial daytime NEE = $5.1 \mu\text{mol m}^{-2} \text{s}^{-1}$, the C flux exponentially decreased at a
303 rate of $\sim 50\%$ of its earlier value, whereas with an initial NEE efflux $\sim 2.5 \mu\text{mol m}^{-2} \text{s}^{-1}$, the C flux decreased at
304 a rate of 100% (Fig. 4). Thus, total C efflux was a contribution of the initial change of NEE and the time taken
305 to return to basal values (i.e., decreasing rates).

306 According to BRT analysis, the factor that most influenced the priming effect in the respiration-dominated
307 period was the change of soil water content at 2.5 cm depth ($\Delta\text{VWC}_{2.5}$; relative importance, RI = 18%), which
308 was followed by the previous NEE (preNEE; RI = 14%), the previous VWC at 2.5 cm depth (RI=14%) and the
309 size of PPT event (RI = 13%). All the other factors had individual RI values lower than 10% (Table 1; Fig. 6).
310 Maximum ΔNEE values were observed at i) larger changes of soil water content at 2.5 cm depth (Fig. 6a), ii)
311 previous neutral NEE (i.e., NEE $\sim 0 \mu\text{mol m}^{-2} \text{s}^{-1}$; Fig. 6b), iii) previous dry soil water content at 2.5 cm depth
312 (Fig. 6c), and iv) with large PPT events ($>15 \text{ mm d}^{-1}$; Fig. 6d). The priming NEE effect decreased farther than
313 these limits. In contrast, in the photosynthesis-dominated period, the previous NEE was the most important
314 factor explaining the “priming effect” (RI=33%). In contrast, the volumetric water content at 15 cm depth, the
315 change of photosynthetic photon flux density, and the volumetric water content at 2.5 cm depth followed in
316 importance (Table 1). Larger changes in NEE (priming effect) were observed at i) more negative previous NEE
317 (i.e., under more photosynthetic activity; Fig. 6e), ii) under drier soil water conditions at 15 cm depth (Fig. 6f),
318 iii) with larger changes of PPFD (decrease of PPFD; Fig. 6g), and iv) under air temperature lower than 16°C
319 and higher than 19°C (Fig. 6h). There was a large interaction between preVWC_{2.5} and PPT for the respiration-
320 dominated period and between preNEE and ΔPPFD for the photosynthesis-dominated period.

321 **3.4 Contribution of priming effect on carbon balance**

322 The carbon balance for these six years for this site was -126 g C m^{-2} , with 2955 and -3080 g m^{-2} of ecosystem
323 respiration and gross ecosystem exchange, respectively, and varied from a sink of $-107 \text{ g C m}^{-2} \text{ y}^{-1}$ to a source
324 of $114 \text{ g C m}^{-2} \text{ y}^{-1}$ (Delgado-Balbuena et al., 2019). Roughly calculation of carbon efflux due to priming effect
325 indicated that extrapolation of mean ΔNEE per event and by year contributes with 142 g m^{-2} for the full six-
326 year period, which corresponds to 5% of total ER flux. In this calculation, parameters like decaying rates, size
327 of PPT event, and previous soil and flux conditions were not considered (modeled) and was subjected to the
328 number of PPT events. Logically, humid years with more PPT events have more contribution of C efflux by
329 priming effect. Each year contributed with less than $30 \text{ g m}^{-2} \text{ y}^{-1}$.

330 **4. Discussion**

331 **4.1 Dynamics of the “Priming effect”**

332 In agreement with the T-D model, NEE exponentially decreased after the PPT pulse (Fig. 5) to almost the pre-
333 PPT NEE rate. The largest C efflux pulses slowly returned to basal C efflux rates and showed larger NEE
334 remnants than the smaller pulses (Fig. 5). This suggests that more persistent VWC quantities achieved with
335 larger size PPT events promoted larger and longer C emissions. If the event was large enough to maintain
336 VWC above a threshold for a long time (e.g., above the wilting point for plants), NEE is expected to remain
337 higher than pre-event rates until nutrients or labile C are depleted (Jarvis et al., 2007; Xu et al., 2004). In
338 contrast, when the PPT event is small, and the soil remains wet for a short time, the C flux peak will be small
339 and less persistent because of soil dry-out and the activity of microorganisms it is likely to end before soil
340 nutrients are depleted. Thus, ‘priming effect’ decaying rates ($-k$) likely are more an issue of water availability
341 than nutrient or C source depletion.

342 **4.2 Thresholds and time delays of the “Priming carbon flux effect”**

343 In our study, the NEE increased immediately (short-time delay) after a PPT event, in accordance with (H1).
344 Moreover, the minimum size of a PPT event needed to detect NEE change was as low as 0.25 mm d^{-1} , in
345 agreement with (H2). We interpret that immediate daytime PPT-induced responses in NEE and ER rates were
346 dominated by heterotrophic respiration and assume that these microbial communities have evolved to take
347 advantage of this short-term water availability. Short-term responses of $< 30\text{-min}$ have also been reported in
348 studies that analyzed soil microorganism activity through molecular and stable isotope techniques (Placella et
349 al., 20012; Unger et al., 2010). Fungi and bacteria on the soil surface have the capability for water-induced re-
350 activation within 1 to 72-h after a PPT event (Placella et al., 2012). Immediate positive NEE increase observed
351 in our study (Fig. A.3) may have resulted from rapid activation of bacteria displaying highest activity 1-h after
352 wetting. Biological soil crusts (BSC) are assemblages of microorganisms forming crusts on the soil and rock
353 surfaces (Belnap, 2003) common in arid lands. At our site, the BSC covers up to 70% of plant interspaces in
354 grazing-excluded conditions and up to 30% in overgrazed sites (Concostrina-Zubiri et al., 2014) with the

355 dominance of actinobacteria (e.g., actinomycetes) and cyanobacteria, which are identified as rapid responders
356 (Bowling et al., 2011).

357 The maximum priming NEE effect was identified under changes larger than 8% of soil water content at 2.5 cm,
358 previous dry soil, neutral previous NEE, and PPT events > 15 mm. These limits may be defined by several
359 conditions, including; 1) the largest and most intense events did not completely infiltrate into the soil, forming
360 abundant runoff and moderating the amount of water penetrating the soil profile at a similar depth as that
361 observed for large-size PPT events, 2) oxygen and CO₂ diffusion limitation under high soil VWC dampened
362 soil respiration, 3) all soil aggregates are disrupted at medium soil VWC likely providing no additional nutrient
363 or C substrate at higher VWCs (Bailey et al., 2019; Lado-Monserrat et al., 2014; Homyak et al., 2018; Chen et
364 al., 2019), and 4) a combination of any of these three. A linear relationship between PPT event size, preVWC_{2.5}
365 and Δ VWC_{2.5} (Fig. 2d) showed that there was not a substantial limitation of water infiltration into the soil at
366 shallow depths, discarding in some way the first condition, whereas the reduction of ER rates in nighttime NEE
367 data after VWC_{2.5} > 12%, and daytime Δ NEE reductions under higher preVWC_{2.5} supports the second
368 mechanism (Fig. 6, and A4).

369 **4.2 The ER and GEE threshold and time delays difference**

370 The smallest PPT events only stimulated ER rates, with no apparent change observed in GEE (Fig. 3). Even a
371 large PPT event of 20 mm d⁻¹ recorded in May 2013 (Fig. 3) did not induce an increase in GEE. In contrast,
372 larger or consecutive PPT events that reached deeper soil profiles stimulated GEE (cumulative PPT > 40mm).
373 These results also explain why the previous soil moisture and the change of soil moisture (2.5 cm depth) better
374 explained Δ NEE at the respiration-dominated period rather than soil moisture at 15 cm depth (Fig 5); this
375 confirms our notion that soil microorganism activity was the source of the immediate CO₂ efflux. In contrast,
376 VWC at 15 cm depth was the second most important factor explaining the priming NEE effect in the
377 photosynthesis-dominated period. Additionally, the change of PPFD during the photosynthesis-dominated
378 period positively affected the priming effect (Fig. 6), meaning that cloudy conditions reduced carbon uptake
379 rather than PPT and stimulated ecosystem respiration.

380 The low PPT threshold that stimulated ER agrees with results from other studies in arid ecosystems (and are
381 even lower). PPT events as small as 3 mm induced respiration of biological soil crusts (Kurc and Small, 2007),
382 and PPT events <10 mm d⁻¹ on a shortgrass steppe promoted net loss of C (Parton et al., 2012). Moreover,
383 Medina-Roldán et al. (2013) at the same study site showed an increase of 36% and 34% of extractable NH₄⁺
384 and NO₃⁻, respectively, after a PPT event of 10 mm, which is an indicative of soil biological activity. However,
385 the dominant species at our site, *B. gracilis*, was reported to respond to PPT events as small as 5 mm (Sala and
386 Lauenroth, 1982), which was the PPT threshold we were expecting. Instead, this study found that large or
387 consecutive PPT events had to occur before an effect on GEE was observed (Fig 3). Nevertheless, we highlight
388 that small PPT events in arid ecosystems that do not lead to C uptake may alleviate stress after severe droughts,
389 rehydrating plant tissues and helping plants to respond faster after larger PPT events (Sala and Lauenroth, 1982;
390 Aguirre-Gutiérrez et al., 2019; Thomey et al 2011).

391 Causes of larger time-delays in GEE than ER are likely due to the delay between the PPT event and the
392 infiltration of water to a given soil layer (e.g., 15 cm depth; Fig. 2e), and the time spent for regrowing of new
393 roots and leaves (Ogle and Reynolds, 2004). These processes promote C losses rather than C uptake in the early
394 growing season (Huxman et al., 2004; Delgado-Balbuena et al., 2019). In contrast, ER was primarily controlled
395 by soil moisture at shallow soil layers that moist immediately after any PPT event and may activate soil
396 microorganism just a few hours after soil wetting as discussed above.

397 **4.3 Influence of event size and a priori conditions**

398 The magnitude of the priming effect was determined by the size of the PPT event and mainly by the ΔVWC as
399 well as the previous condition of the ecosystem (i.e., previous C flux and previous soil VWC). These results
400 agree with (H3) that proposed the PPT event size and previous conditions of the semiarid grassland would
401 control the magnitude of the “priming NEE effect”. The previous VWC offers insight into the potential dry-
402 wet shock experienced by soil aggregates and microorganisms (Haynes and Swift, 1990) and thus accounts for
403 nutrient and labile C accumulation in soil (Bailey et al., 2019).

404 Results indicated that larger C effluxes were induced from medium amount of PPT when the previous soil
405 conditions were dry and had an initial value of $NEE = \sim 0$. Several mechanisms can explain this result: i) the
406 accumulation of nutrients and labile C into the soil (Schimel and Bennet, 2004) because low activity of
407 microorganisms ($NEE \sim 0$) under dry soil (Homyak et al., 2018), ii) if soil VWC is maintained for an extended
408 period above a threshold, then soil microbial activity exhaust labile C sources (Jarvis et al., 2007; Fierer and
409 Schimel, 2002). Consequently, recalcitrant C sources subjected to microbial decomposition decrease
410 mineralization rates (Van Gestel et al., 1993).

411 **4.4 Importance of the priming effect in the annual C balance.**

412 Our results do not support the hypothesis that a significant contribution of C release from the “priming effect”
413 decreases the net annual C uptake of the semiarid grassland (H4). The contribution of these short-term C efflux
414 events to annual C balances accounted for a considerable amount, but it was a small contribution if it is
415 considered into the ecosystem respiration flux, which was almost $3000 \text{ g m}^{-2} \text{ s}^{-1}$ (Delgado-Balbuena et al., 2019).
416 Notwithstanding its contribution is low ($\sim 5\%$ of ecosystem respiration), it is important considering that the
417 annual C balance (NEE) is a small fraction of the difference between ER and GEE. Thus, a 5% of C released
418 represents up to 500% of the net C uptake during an almost neutral year and may turn a C sink ecosystem into
419 a net C source.

420 **4.5 Priming effect and climate change perspectives.**

421 The low $\Delta SWC_{2.5}$ and PPT threshold for respiration suggests that almost all PPT events occurring in the
422 semiarid grasslands will produce C efflux but will be limited by the characteristics of the PPT pattern and
423 previous soil conditions at the site. Therefore, we expect that small PPT events with previous dry conditions
424 or long inter-event periods will limit the priming effect by maintaining the system below threshold conditions.

425 Moreover, consecutive PPT events or large PPT events should keep soil water content above a threshold that
426 will promote C uptake by photosynthesis, which in the long term will overcome C loss from the priming effect.
427 However, climate change scenarios forecast for the semiarid grassland in Mexico a decrease in winter PPT and
428 the increase in storms with larger inter-event periods, which are conditions for increasing the amount of C
429 released by the priming effect (Arca et al., 2021; Darenova et al., 2019).
430 Further analysis of the effect of these PPT events on vegetation is necessary since productivity will also depend
431 on PPT event size and will be modulated by previous soil conditions. Additionally, it is likely that productivity
432 will benefit more from accumulated PPT than respiration. Still, more analysis of projected PPT scenarios is
433 required to accurately forecast the contribution of the Birch effect to C balance under more frequent droughts.
434 In this sense, parameterizing a model like the T-D model will provide valuable information on more accurate C
435 effluxes from the priming effect and how it will be affected by changes in precipitation pattern. Only after that
436 will we be able to predict the course of the semiarid grassland as a source or sink of C under PPT pattern
437 changes.

438 **5. Conclusions**

439 Previous soil water conditions and previous NEE were the most important factors controlling the priming effect
440 in the semiarid grassland. The precipitation size had an important role in explaining the priming effect but only
441 in the respiration-dominated period. Delays between change responses at deeper soil layer and for regrowing
442 processes could hide the relationship between precipitation and priming effect during the photosynthesis-
443 dominated period. The importance of the priming effect in the carbon balance could be more relevant under
444 forecasted changes in precipitation patterns by increasing in both the frequency and intensity the dry-wet soil
445 cycles. Further analysis of the effect of this change of precipitation pattern on ecosystem productivity is
446 necessary before we can conclude about changes in the carbon balance of the semiarid grassland.

447

448 *Author contributions.* The study was conceived by JD, TA, HL, and RV. JD, TA and CAA get and processed
449 eddy covariance data. JD, TAR, and LFM implemented the method and performed the data analyses. TAR and
450 CAA get and processed the Enhanced Vegetation Index data. TA, HL, LFM, and RV helped to interpret the
451 results. JD, TA, HL, and RV prepared the first draft, and all authors contributed to discussion of results and the
452 revisions of the paper.

453

454 *Competing interests.* The authors declare that they have no conflict of interest.

455

456 *Availability of data.* The datasets used for this study are available at Zenodo
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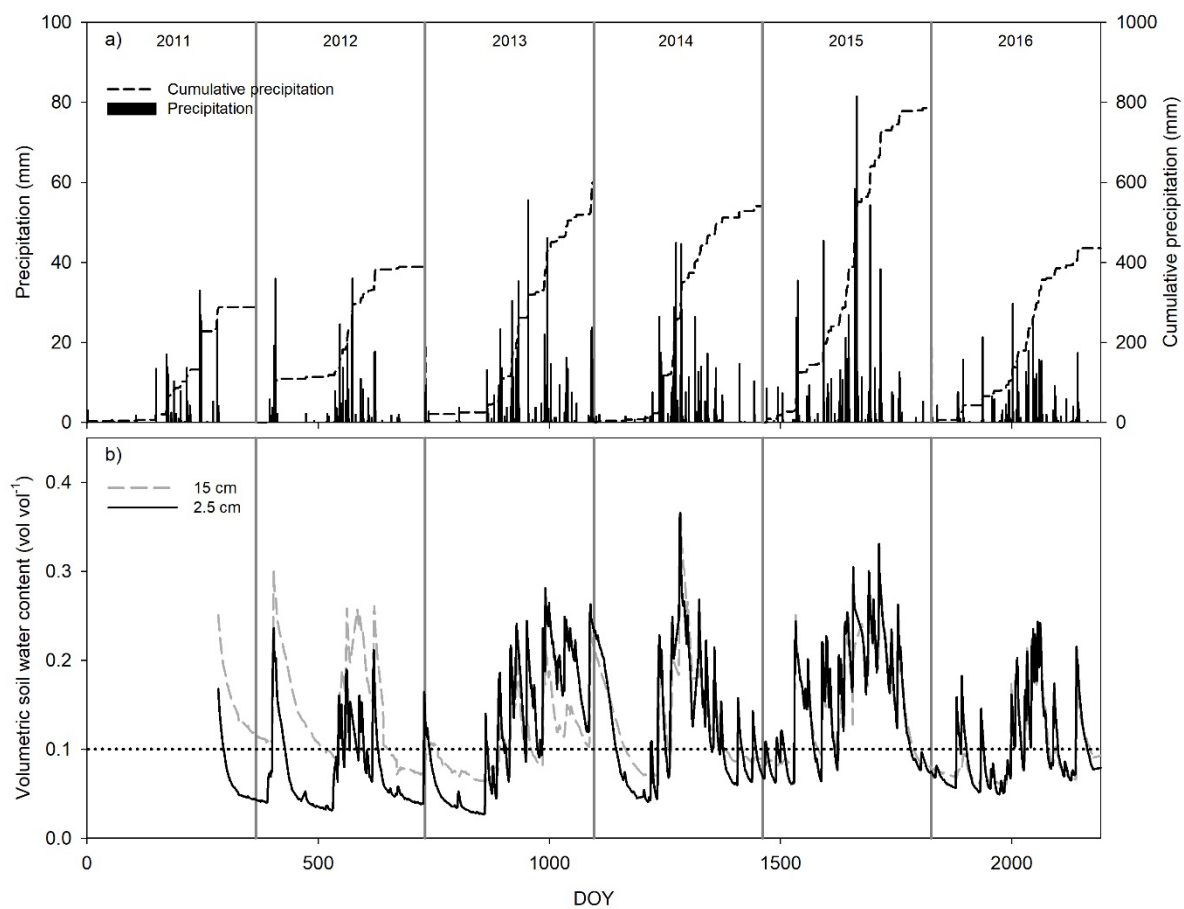
635 **Table 1.** Relative importance (RI) of the first four most important environmental factors for the “priming CO₂ effect”.

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Respiration-dominated period	RI
$\Delta\text{VWC}_{2.5}$	18.66
preNEE	14.67
preVWC _{2.5}	14.08
PPT	13.64
preVWC ₁₅	8.09
VWC _{2.5}	7.46
Photosynthesis-dominated period	
preNEE	33.32
VWC ₁₅	12.25
ΔPPFD	11.52
VWC _{2.5}	9.16
Tair	8.32
preVWC _{2.5}	7.79
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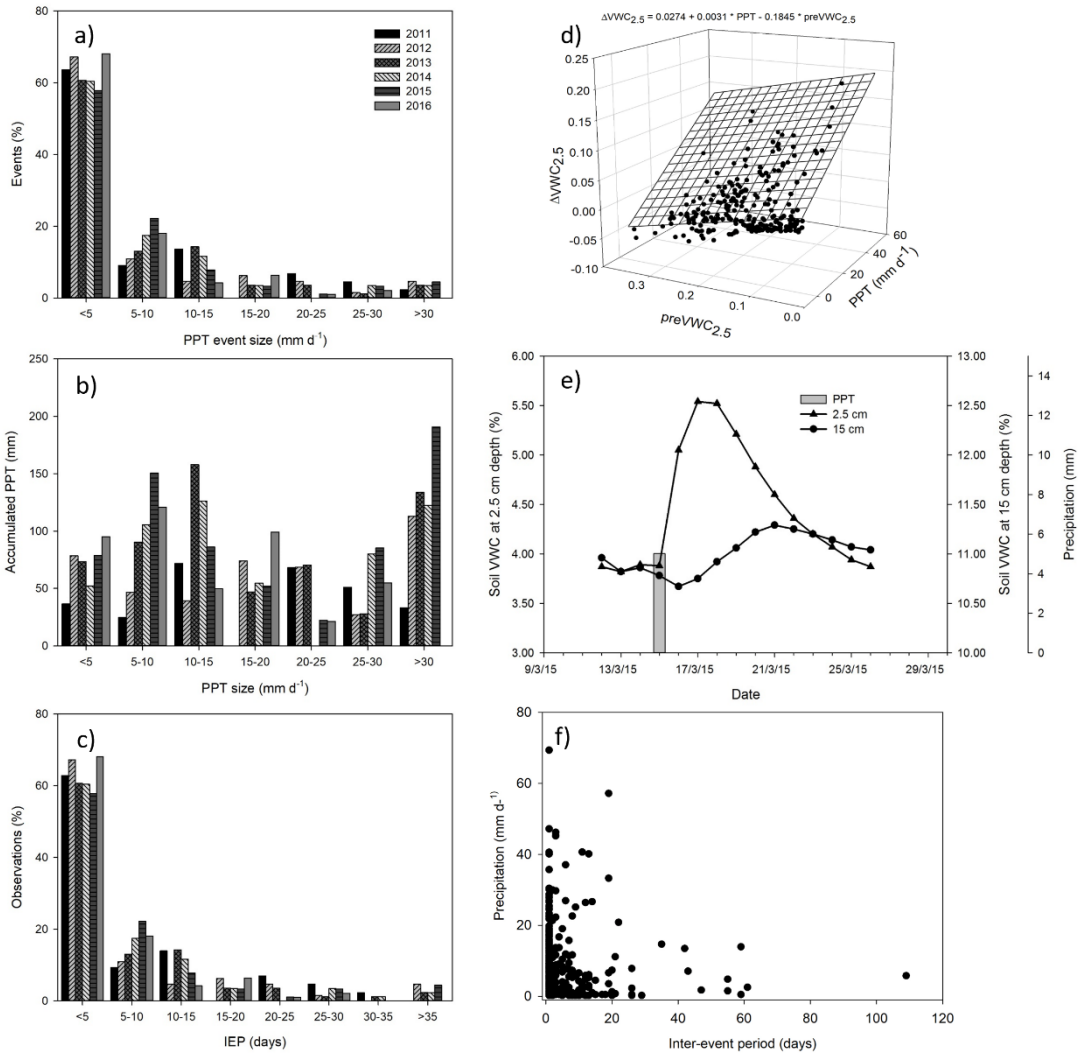
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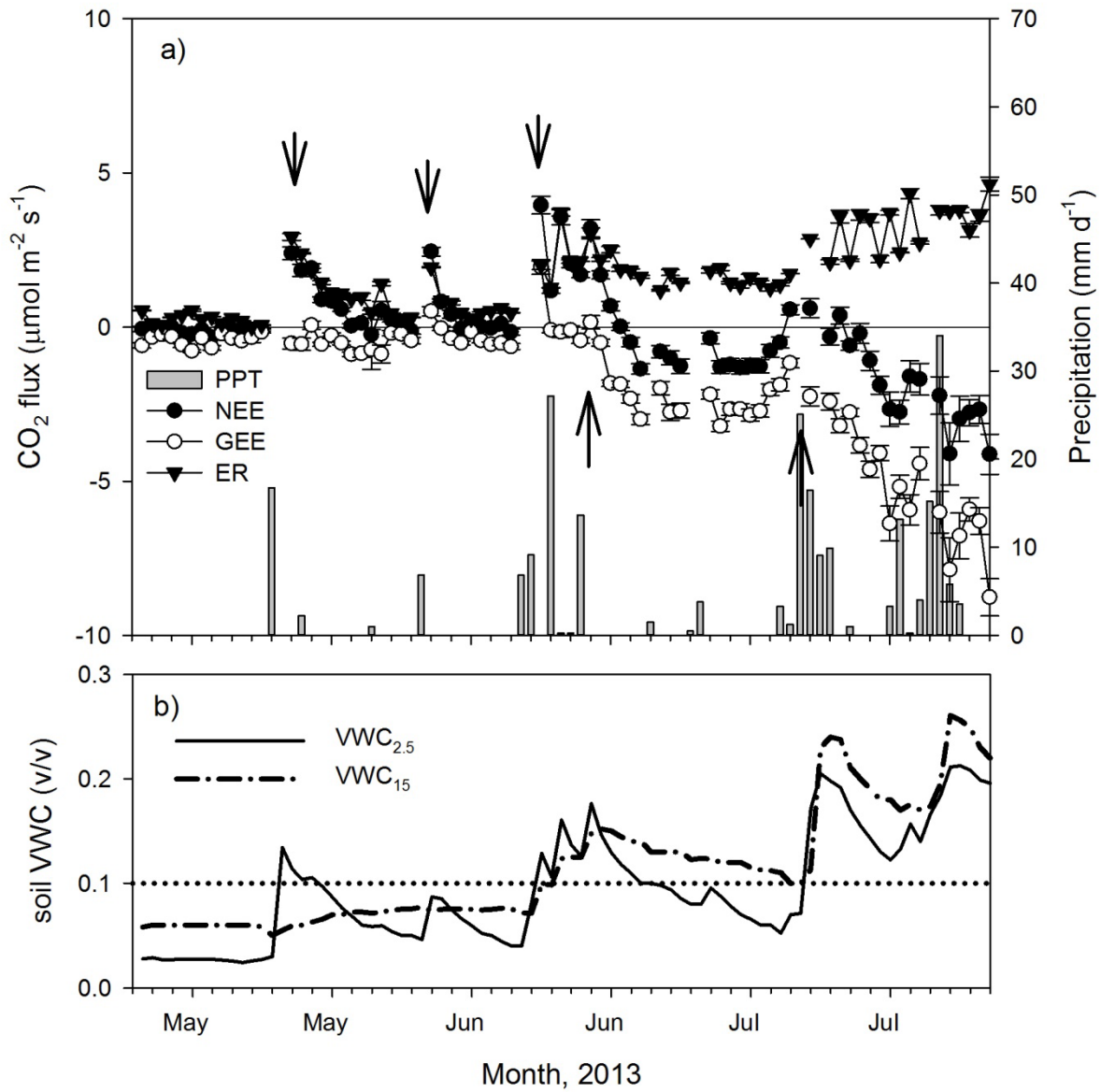
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640 **Figure 1.** Seasonal and interannual variation of daily precipitation and cumulative precipitation (a), and volumetric soil
 641 water content at 2.5 (black line) and 15 cm depth (gray line; b). Dotted line at 10% of soil was content was depicted as
 642 reference.



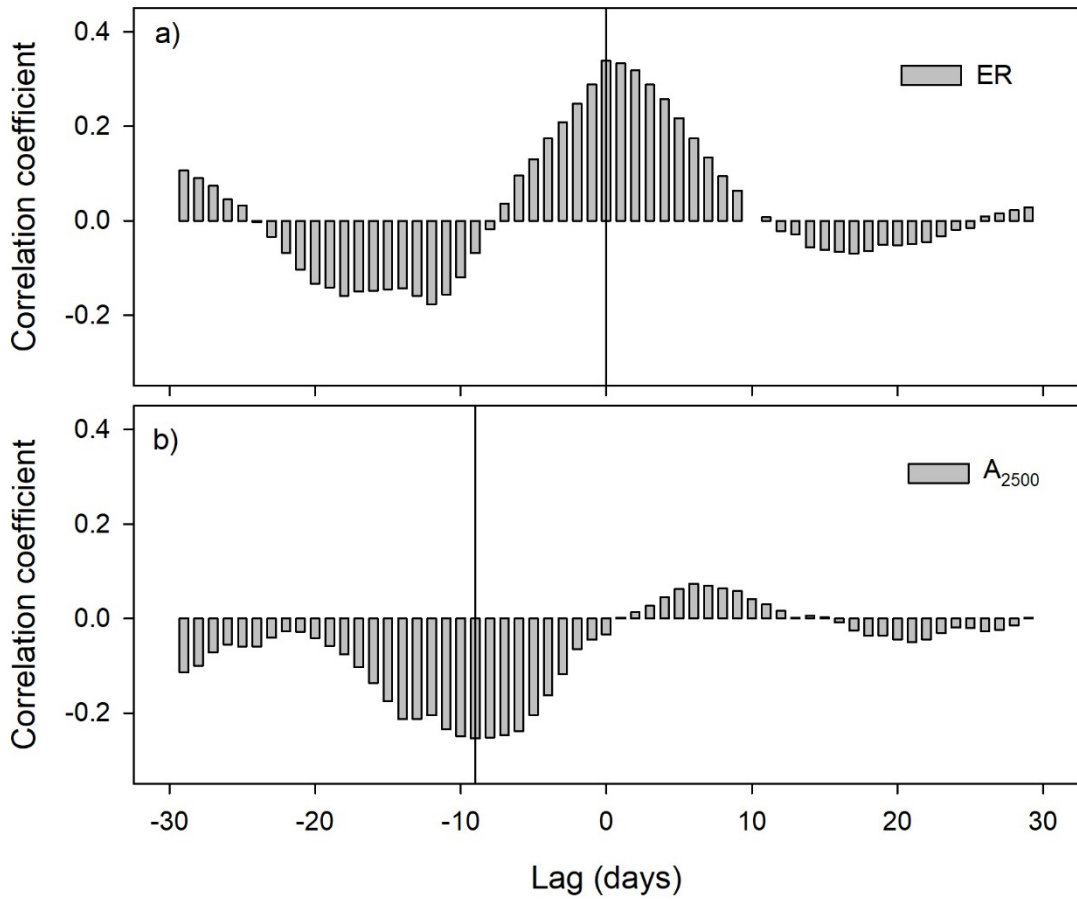
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644 **Figure 2.** Characterization of precipitation pattern. Histogram of the size of precipitation events through six years (a), the
 645 accumulated precipitation by size of precipitation event (b), and the number (%) of precipitation events by inter-event period
 646 classes (IEP, days; c). Relationship between size of precipitation event (mm d^{-1}), previous volumetric soil water content at
 647 2.5 cm depth (v/v) and the change in soil volumetric water content at 2.5 cm depth (v/v). Dynamic of soil water content at
 648 two depths (2.5 and 15 cm) after a precipitation event of 5 mm through the time (e), and relationship between inter-event
 649 period and the size of precipitation event (f).



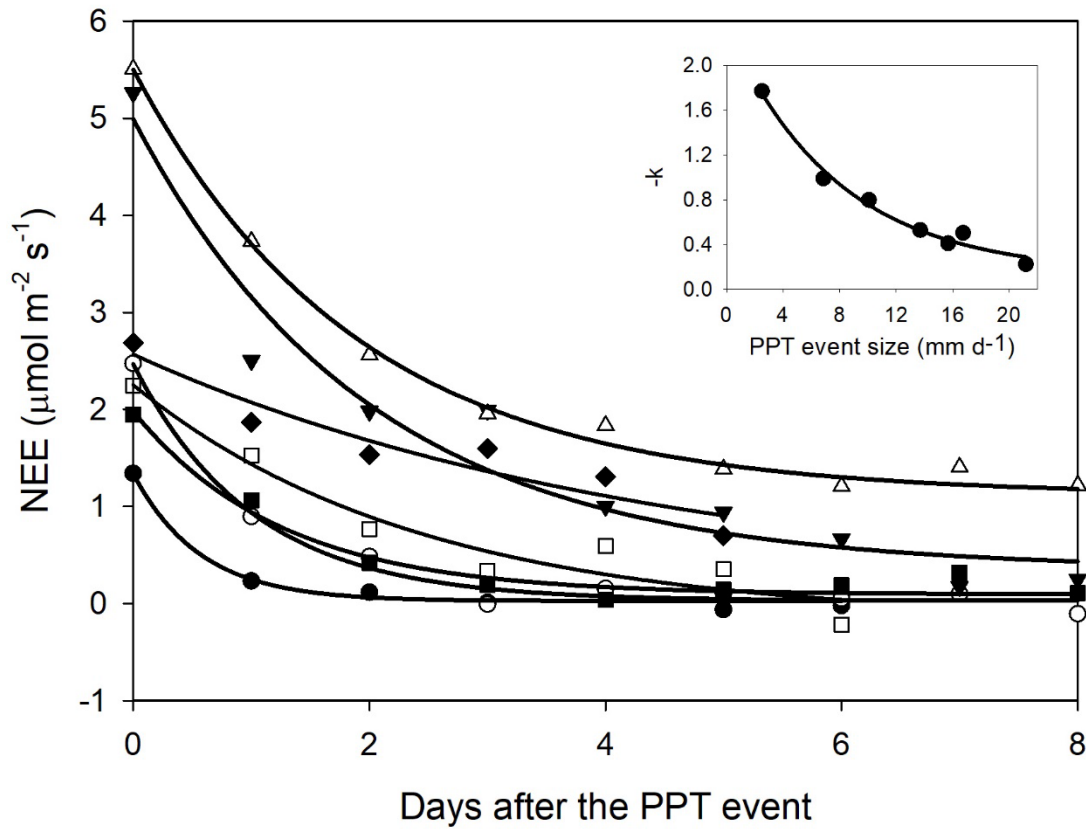
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651 **Figure 3.** Dynamics of a) precipitation (mm d^{-1}) and net ecosystem exchange (NEE, $\mu\text{mol m}^{-2} \text{s}^{-1}$, daily means ± 1 SE) and
 652 its components, the gross ecosystem exchange (GEE, $\mu\text{mol m}^{-2} \text{s}^{-1}$) and ecosystem respiration (ER, $\mu\text{mol m}^{-2} \text{s}^{-1}$) for the
 653 transition from the dry (December – May) to the wet season (June – November) in 2013. b) volumetric soil water content
 654 dynamics (VWC, v/v) at two depths (2.5 cm and 15 cm). Arrows indicate apparent changes in GEE and ER trends. The
 655 dotted line indicates $\text{SWC} = 0.1$.



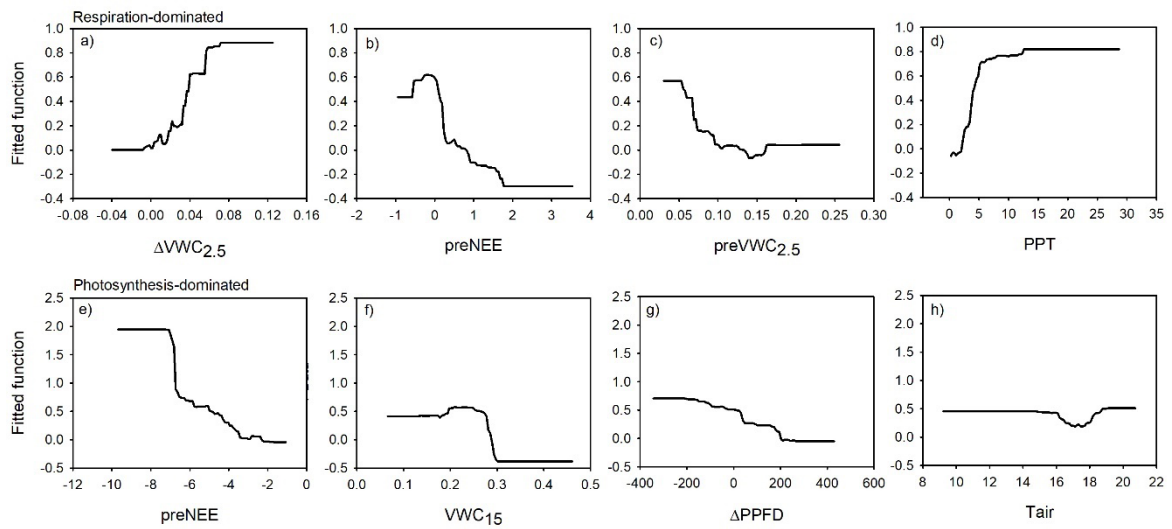
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657 **Figure 4.** Cross-correlation coefficients between detrended time series of soil water content at 2.5 cm depth and ecosystem
 658 respiration (ER, a), and between soil water content at 15 cm depth and photosynthesis at 2500 $\mu\text{mol m}^{-2} \text{s}^{-1}$ of photosynthetic
 659 photon flux density (A_{2500} ; b).



660

661 **Figure 5.** Net ecosystem exchange (NEE) after a precipitation event showing the decreasing effect through time (days).
 662 The decreasing effect rate was adjusted to an exponential negative model $NEE = y_0 + a \cdot \exp(-k \cdot t)$. The insert stands for
 663 the relationship between the decaying rate ($-k$) and the PPT event that originated the NEE change. This relationship was
 664 fitted with an exponential model (black line; $-k = y_0 + a \cdot \exp(-b \cdot \text{PPT_event})$). Symbols indicate different PPT event sizes
 665 that originated the NEE change, 13.7mm d⁻¹ (Δ), 16.74 mm d⁻¹ (▼), 6.86 mm d⁻¹ (○), 10.08 mm d⁻¹ (■), and 2.52 mm d⁻¹
 666 (●).



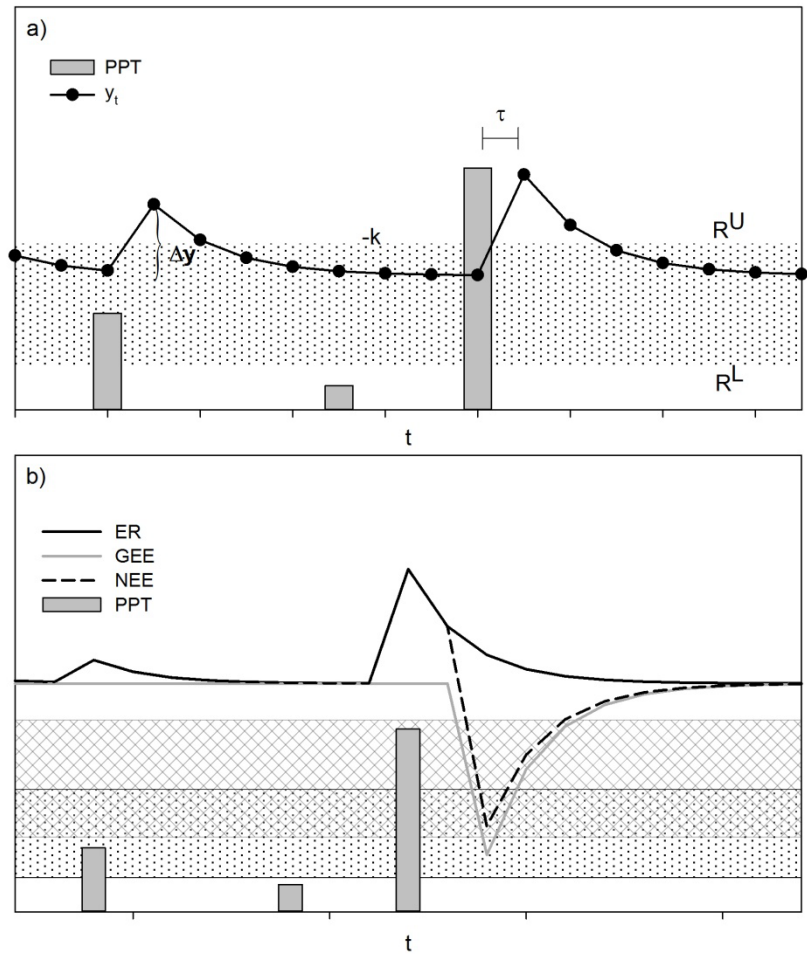
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668 **Figure 6.** Fitted functions of the boosted regression trees between the “priming CO₂ effect” and the four most important
 669 environmental variables at ecosystem respiration-dominated period (upper panel) and at the photosynthesis-dominated
 670 period (bottom panel). Priming effect (ΔNEE , $\mu\text{mol m}^{-2} \text{s}^{-1}$); previous NEE (preNEE, $\mu\text{mol m}^{-2} \text{s}^{-1}$); previous VWC at 2.5
 671 cm depth (preVWC_{2.5}, v/v); PPT event size (PPT, mm); VWC at 15 cm depth (VWC₁₅, v/v); change of photosynthetic
 672 photon flux density ($\Delta PPFD$, $\mu\text{mol m}^{-2} \text{s}^{-1}$); air temperature (T_{air}, °C).

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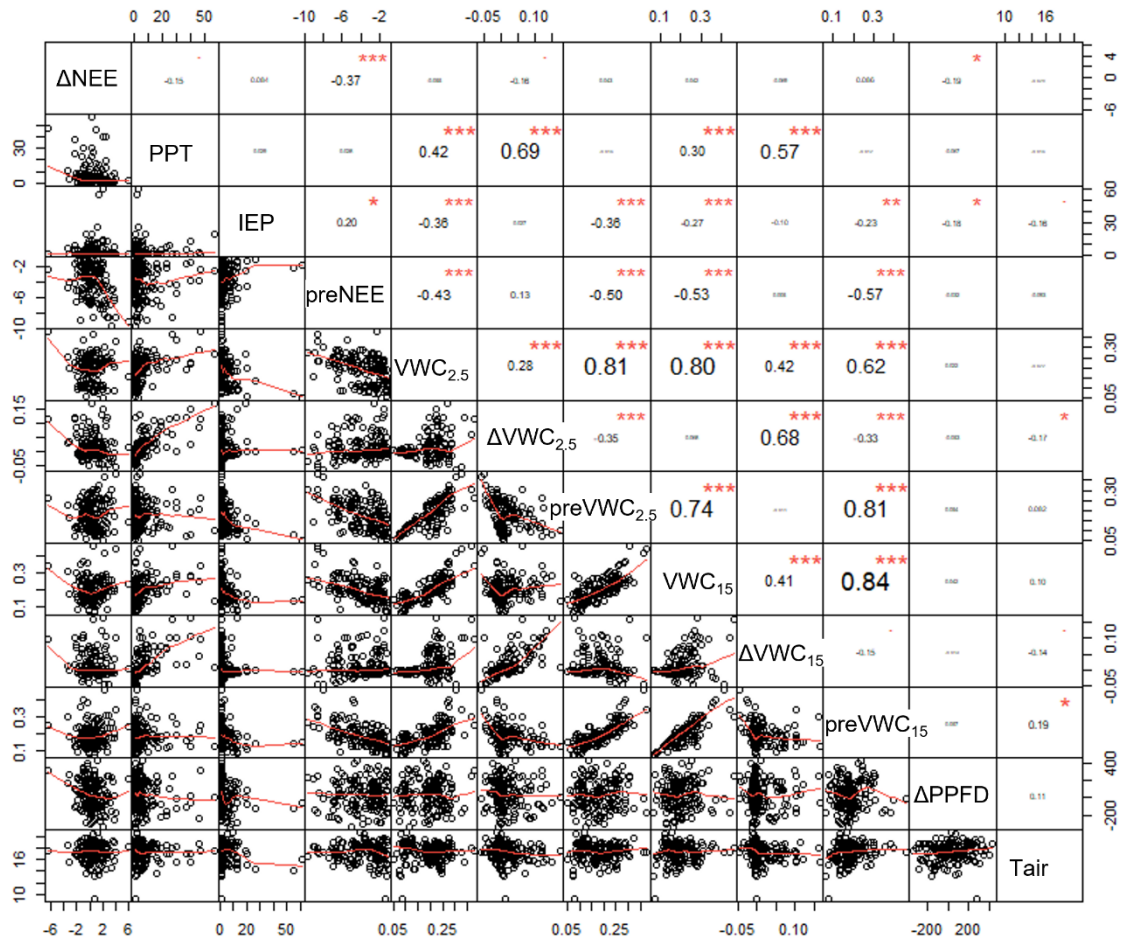
675 **Appendix A. Ancillary figures**



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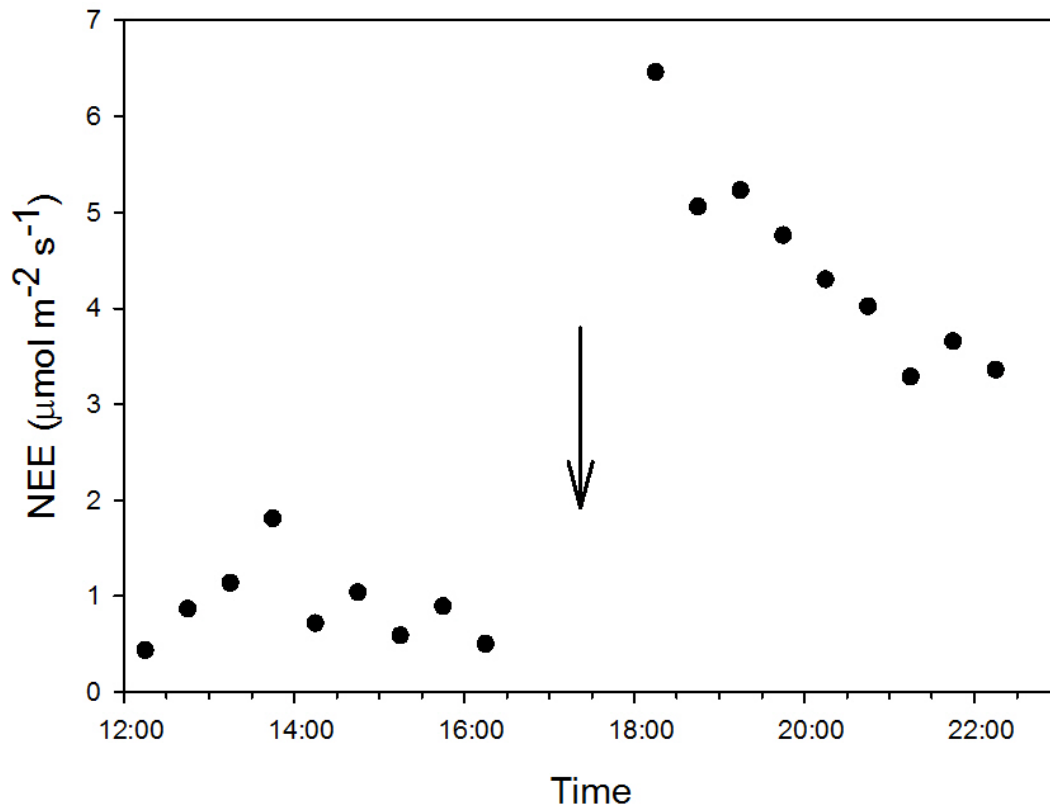
677 **Figure A1.** The Threshold-Delay model (Ogle and Reynolds, 2004). a) The magnitude of the increase in the response
 678 variable (Δt , e.g. carbon flux, y_t) is determined by the size of PPT event and by the previous state of the response variable.
 679 The decreasing rate of the response following the stimulus is denoted by $-k$. The low PPT threshold (R^L) indicates the
 680 minimum size PPT event to stimulate a response, and the upper PPT threshold (R^U) indicates PPT events that do not cause
 681 additional increment in the response variable. The time interval between the stimulus and the response is described by τ . b)
 682 The response of the net ecosystem exchange (NEE), that is the balance between the gross ecosystem exchange (GEE) and
 683 ecosystem respiration (ER), vary in response to changes of GEE and ER. According to the T-D model, GEE and ER have
 684 different PPT thresholds (dotted band and mesh stand for effective PPT events size for ER and GEE, respectively), with ER
 685 responding to smaller size PPT events than GEE, therefore, small PPT events favor C release whereas large PPT events
 686 stimulate net C uptake by the ecosystem. Differences of time responses between soil microorganisms and plants to soil wet

687 up led GEE and ER to differ in time delays (τ), with shorter time delays for ER than GEE (Huxman et al., 2004a). The
688 hypothetical curve for NEE and its components was calculated introducing arbitrary parameters in the T-D model equations
689 of Ogle and Reynolds (2004).



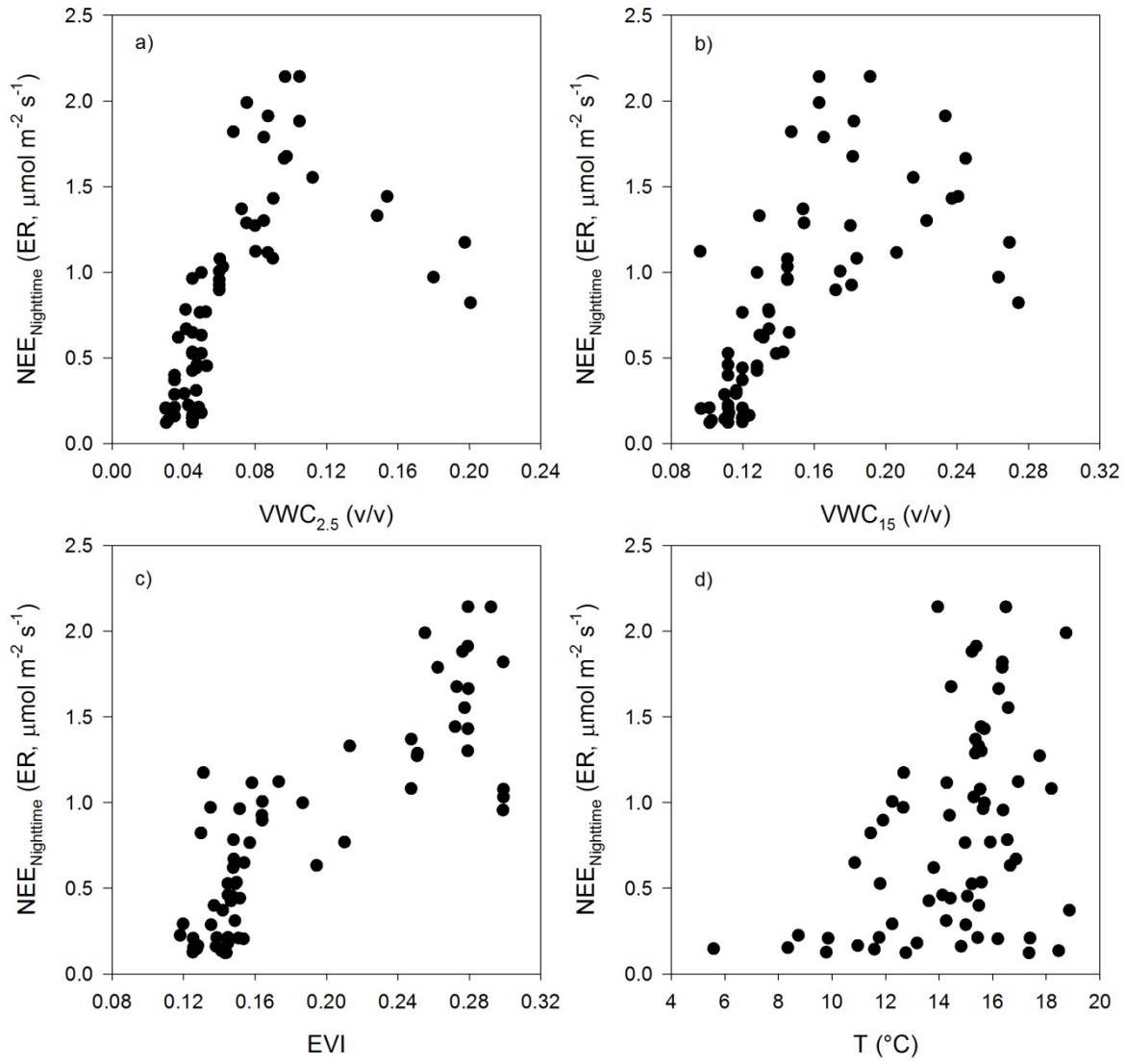
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691 **Figure A2.** Correlation matrix among all variables.



692

693 **Figure A3.** Dynamic of half an hour net ecosystem exchange ($\mu\text{mol m}^{-2} \text{s}^{-1}$) after a precipitation event of 8.12
 694 mm. The arrow indicates the time of PPT event occurrence.



695

696 **Figure A4.** Relationship between nighttime-NEE derived ER and a) the soil volumetric water content at 2.5 cm
 697 depth ($VWC_{2.5}$, v/v), b) the soil volumetric water content at 15 cm depth (VWC_{15} , v/v), c) the enhanced
 698 vegetation index (EVI), and d) the air temperature (T , $^\circ\text{C}$).