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10	Amazonian Biogenic Volatile Organic Compounds under Global Change
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29 ecological interactions, depositional processes, land use, land cover, global change

30

31 Abstract

32 Biogenic volatile organic compounds (BVOCs) play important roles at cellular, foliar, ecosystem, and atmospheric 33 levels. The Amazonian rainforest represents one of the major global sources of BVOCs, so its study is essential for 34 understanding BVOC dynamics. It also provides insights into the role of such large and biodiverse forest ecosystem in 35 regional and global atmospheric chemistry and climate. We review the current information on Amazonian BVOCs 36 and identify future research priorities exploring biogenic emissions and drivers, ecological interactions, atmospheric 37 impacts, depositional processes, and modifications to BVOC dynamics due to changes in climate and land cover. A 38 feedback loop between Amazonian BVOCs and the trends of climate and land-use changes in Amazonia is then 39 constructed. Satellite observations and model simulation time series demonstrate the validity of the proposed loop 40 showing a combined effect of climate change and deforestation on BVOC emission in Amazonia. A decreasing trend 41 of isoprene during the wet season, most likely due to forest biomass loss, and an increasing trend of the sesquiterpene 42 to isoprene ratio during the dry season, suggest increasing temperature stress induced emissions due to climate 43 change.

44

45 **1. Introduction**

Biogenic volatile organic compounds (BVOCs) are emitted by vegetation, bacteria, algae, fungi, and 46 47 animals. Their rates of emission vary greatly in response to biotic (e.g. herbivory) and abiotic (meteorological) factors (Šimpraga et al., 2019). They have diverse roles at multiple scales, from cellular 48 49 protection and defence at the foliar level, through chemical signalling at regional level, up to influencing 50 rainfall at the ecosystems scale (Laothawornkitkul et al., 2009). BVOCs profoundly affect biosphere-51 atmosphere interactions by atmospheric reactivity, aerosol growth processes, and cloud formation, and 52 therefore radiative balance (Kulmala et al., 2013; Nölscher et al., 2016; Pöschl et al., 2010). As BVOCs are 53 precursors of aerosols, they can impact the regional radiative forcing directly through aerosol scattering and 54 indirectly by influencing cloud processes. Furthermore, higher cloud cover leads into increased diffuse radiation, boosting net primary productivity and thereby BVOC emissions (Rap et al., 2018). However, 55 56 BVOCs can have further impacts on the radiative forcing by their influence on tropospheric ozone 57 formation or by increasing the lifetime of methane by suppression of its major sink, the OH radical 58 (Peñuelas & Llusià, 2003).

59 Vegetation is the largest source of BVOC emissions, with tropical trees responsible for ca. 80% of global 60 terpenoid emissions and 50% of other BVOC emissions (Guenther et al., 2012). With half of the global 61 tropical forests, Amazonia is the largest remaining pristine continental area (Andreae et al., 2015; Pöhlker et al., 2019). Amazonia stores 50% of tropical-forest carbon, plays a vital role in the global cycles of 62 energy, water, and carbon, and has the largest biodiversity in the world (Levine et al., 2016; Malhi et al., 63 2008; Swift et al., 1998; ter Steege et al., 2013). This dynamic system emits large quantities of chemically 64 65 diverse BVOCs to the atmosphere, with implications both within and above Amazonian forest canopies. 66 Amazonia is highly sensitive to changes in climate and land use due to its strong energy and biochemical 67 cycling (Levine et al., 2016). It has been established that recent human action is already altering Amazonian conditions and feedbacks (Sena et al., 2018; Shrivastava et al., 2019), complicating the 68 69 understanding of BVOCs in this region. In fact, BVOC emission decrease due to forest conversion to 70 croplands in the tropics has been shown to result in a positive radiative forcing (Jia et al., 2019; Scott et al., 71 2018). Additionally, the impact of Amazonian BVOCs on global primary production through diffuse 72 radiation has large implications for the global carbon balance (Rap et al., 2018). Furthermore, by 73 participating in cloud formation processes, Amazonian BVOCs play a key role in the regional hydrological 74 cycle, since Amazonian rainforests contribute to precipitation throughout the South American continent 75 (Boers et al., 2017; Rocha et al., 2018). Therefore, it is critical to understand the BVOC response to 76 environmental stresses caused by changes in climate and land use in Amazonia, and how these changes will 77 feedback into the Earth system (Peñuelas & Staudt, 2010).

We here review the recent advances in Amazonian BVOC research and estimate potential future BVOC
dynamics using information currently available for emission processes, resulting ecological interactions
and atmospheric impacts, depositional processes, and effects and feedbacks of climate and land use change.
We propose a feedback loop, which is supported by satellite observations and model simulations of BVOC
fluxes, to identify research priorities in Amazonian BVOCs.

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2. Advances on BVOC scientific research in Amazonia

a. History of measurements

Measurements of BVOCs in Amazonian ecosystems are logistically difficult due to accessibility, and power constrains. Research on this topic began in the 1970s and 1980s, when tethered balloon observations demonstrated that Amazonia is a major source of BVOC emissions to the atmosphere (Crutzen et al., 1985; Greenberg & Zimmerman, 1984; Zimmerman et al., 1988) providing understanding into atmospheric chemistry (Grosjean, 1995). The establishment of the Large-Scale Biosphere-Atmosphere Experiment in Amazonia (LBA) provided the first large-scale estimates of isoprene emissions and oxidation products from data collected by aircraft and tower-based stations (Kesselmeier et al., 2009). Nevertheless,

92 understanding of the controlling processes was still scarce due to the lack of data on temporal and spatial 93 variability and the very limited number of BVOCs species studied (Kesselmeier et al., 2009). The most 94 commonly used prominent technique in early studies was BVOC collection in cartridges, followed by transportation of field samples to laboratories outside of Amazonia for analysis by gas chromatography. 95 96 The logistical difficulties for sample transport were addressed by establishing more locations for tower-97 based measurements, long term field campaigns, new sampling approaches such as the use of drones 98 (McKinney et al., 2019), along with more advanced analytical tools such as proton-transfer-reaction mass 99 spectrometry (Bracho-Nunez et al., 2013; Crutzen et al., 2000; Fuentes et al., 2016; Jardine et al., 2011b, 100 2011a, 2011c, 2014, 2016b; Karl et al., 2009; Kesselmeier et al., 2009; Kuhn et al., 2007; Liu et al., 2016; Rinne et al., 2002; Williams et al., 2001; Yáñez-Serrano et al., 2015), the comparative reactivity method 101 102 (CRM) (Nölscher et al., 2016; Pfannerstill et al., 2018; Sinha et al., 2008), photoionization detector (PID) 103 (Harley et al., 2004), semi-volatile thermal desorption aerosol gas chromatography (SV-TAG) (Yee et al., 104 2018), and isoprene portable monitoring system (T. Taylor personal communication). Extensive campaigns 105 and projects such as GoAmazon and ATTO provided integrated investigations of emissions, chemistry, 106 transport, and cloud processes (Andreae et al., 2015; Martin et al., 2017). Seasonality was clearly addressed 107 (Alves et al., 2016, 2018; Nölscher et al., 2016; Yáñez-Serrano et al., 2015, 2018b) and greater effort was 108 expended towards obtaining a mechanistic understanding that was provided by foliar-level measurements 109 under natural and controlled experiments (Alves et al., 2014; Bourtsoukidis et al., 2018; Bracho-Nunez et 110 al., 2012, 2013). Despite the increasing number of observation datasets and efforts toward addressing horizontal heterogeneity (Batista et al., 2019), comparing ecosystems remains difficult due to the spatial 111 112 diversity of Amazonia (see S.I.1) and the inadequate information over controlling processes. Figure 1 shows the number and location of BVOC measurements available since 1984, when the first measurements 113 were published, showing the lack of spatial variability in the Amazonian BVOC studies to date. 114

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b. History of modelling

There are still too few observations of BVOC emissions for accurate representation of the regional 116 117 variability, resulting in large uncertainty in model predictions. Initially, the BVOC emissions algorithm 118 proposed by Guenther et al., (1995) included light- and temperature-dependent emissions of isoprene, but 119 all other emissions were regarded as only temperature-dependent, a potentially misleading assumption 120 (Kesselmeier & Staudt, 1999). The algorithm was updated in 1999 by a simple leaf-age activity factor to determine the dependence on meteorological conditions. The Model of Emissions of Gases and Aerosols 121 122 from Nature (MEGAN) then simulated net emission rates, accounting for losses of primary emissions 123 (Guenther et al., 2006), with the latest update adding a longer list of compounds and their emission factors 124 (Guenther et al., 2012; Sindelarova et al., 2014). Further BVOC emissions modelling included Organizing 125 Carbon and Hydrology in Dynamic EcosystEm (ORCHIDEE) (Messina et al., 2016) and Lund-PotsdamJena-General Ecosystem Simulator (LPJ-GUESS) (Acosta Navarro et al., 2014). Amazonian modelling has
thus been improved, but a reliable predictive framework for emission capacities in Amazonia requires an
improved mechanistic understanding of BVOC variation across plant populations and of their interactions
with other living organisms and abiotic factors (Alves et al., 2018; Kuhn et al., 2004a; Salazar et al., 2018;
Taylor et al., 2018).

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c. Satellite applications

132 Amazonian isoprene fluxes can be derived at regional, or global scales via satellite measurements (Barkley 133 et al., 2009, 2013; Bauwens et al., 2016; Fu et al., 2019; Gu et al., 2017). This can improve our knowledge 134 of the dynamics in scale and time. Formaldehyde is formed during the oxidation of most VOCs released in 135 the atmosphere. It is a major intermediate product in isoprene oxidation, and therefore satellite observations 136 of formaldehyde (HCHO) have been used to infer isoprene emissions at global and regional scales (Millet 137 et al., 2008; Pamler et al., 2003; Stavrakou et al., 2009). The variability of emissions estimated with these 138 satellite data generally lies within the variability of the surface observations. Direct observations of 139 isoprene recently acquired by the satellite-borne Cross-track Infrared Sounder (CrIS) (Fu et al., 2019) will open 140 new avenues in the isoprene emission estimation from satellites. A global study using these data found a regional reduction (-2% y⁻¹) of biogenic emissions over Amazonia during the period of 2005 to 2013 but no 141 142 plausible cause could be identified (Bauwens et al., 2016).

143 3. Emission

144 The BVOCs in the Amazonian rainforest have many sources, with vegetation being the most prominent 145 (Kesselmeier et al., 2009) (Figure 2). A large variety of BVOCs are emitted from Amazonian ecosystems. 146 Isoprene is the most strongly emitted compound, with reported ambient mixing ratios of 0.5-15 ppb and 147 fluxes of 0.7-12.1 mg m⁻²_{ground} h⁻¹, although other isoprenoids have been reported such as monoterpenes 148 (with mixing ratios typically below 1 ppb but in some cases up to 5.5 ppb (Greenberg & Zimmerman, 149 1984; Kesselmeier et al., 2002b)), sesquiterpenes (SQT) (with mixing ratios up to 0.16 ppb), and even diterpenes (0.0002 ppb) (Yee et al., 2018). Methanol is the second most emitted compound, with measured 150 151 mixing ratios reported up to 6 ppb (Kesselmeier et al., 2002b; Yáñez-Serrano et al., 2015). Other 152 oxygenated compounds have also been reported, such as acetaldehyde, acetone, methyl ethyl ketone (MEK) 153 (Bracho-Nunez et al., 2012; Rottenberger et al., 2008; Yáñez-Serrano et al., 2016), ethanol (Bracho-Nunez 154 et al., 2012; Rottenberger et al., 2008), formaldehyde (Karl et al., 2009; Kesselmeier et al., 2000), acetic 155 and formic acid (Jardine et al., 2011b; Kuhn et al., 2002a), green-leaf volatiles (GLVs) (Jardine et al., 156 2015b), isoprene oxidation products (including methyl vinyl ketone (MVK), methacrolein (MACR), 157 isoprene hydroperoxide (ISOPOOH), and hydroxyacetone (Karl et al., 2009), aromatics (Paralovo et al., 158 2015), sulphurous compounds such as dimethyl sulphide (Jardine et al., 2014) as well as chlorine compounds such as methyl chloride and chloroform (Gebhardt et al., 2008; Gregory et al., 1986; Yokouchi
et al., 2002) (Table 1, figure 3 and 4). Additionally, the emission, and consequent atmospheric abundance is
given as the sum of enantiomers due to analytical constrains, yet BVOCs are generated by enzymes and are
therefore chiral. Although chemically identical, they can induce differing biological responses. Thus
examining the different enantiomers would lead to a doubling of information (Williams et al., 2007).

164 Vegetation emits BVOCs, as *de novo* compounds from recently photosynthesised carbon or as volatilised compounds from storage pools or resins and waxes (Kesselmeier & Staudt, 1999). These emissions may 165 166 represent a substantial loss of carbon previously assimilated by photosynthesis (up to 2% of carbon fixed for tropical forests) (Kesselmeier et al., 2002a). Isoprene is the most common de novo biosynthesised 167 compound, with isoprene-emitting plants found as a typical component in tropical forests (Harley et al., 168 169 2004; Šimpraga et al., 2019; Taylor et al., 2018). Further species, such as monoterpenes are also released de 170 novo from leaves, glands, and resin ducts (Kesselmeier & Staudt, 1999) in Amazonia (Jardine et al., 2015a; 171 Kuhn et al., 2002b; Piva et al., 2019; Yáñez-Serrano et al., 2018b).

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In addition to living vegetation, leaf litter is an important source of BVOCs (Greenberg et al., 2012), particularly during the start of the dry season when litterfall peaks, and decomposition is further aided by wetness with the upcoming wet season. Little is known about this BVOC source in Amazonia, but monoterpene addition to soils from leaf fall and excreted from plant resins and roots have been reported elsewhere (Delory et al., 2016; Marmulla & Harder, 2014), suggesting a potential, unquantified role of Amazonian ecosystems with a rapid turnover of biomass.

Soils are a poorly explored BVOC source in the Amazonian rainforest, including their microorganisms, roots, and surface reactions (Tang et al., 2019). SQTs have recently been found to be strongly emitted by soil bacteria (Bourtsoukidis et al., 2018), with responses to soil conditions, because microorganisms are highly spatially heterogeneous (Buscardo et al., 2018), dynamic and able to produce and consume BVOCs strongly interacting within the soil ecosystem. Also reduced sulphur compounds such as dimethyl sulphide, carbon disulphide or carbonyl sulphide can be released by soils (Bunk et al., 2017), including Amazonian soils (Jardine et al., 2014).

Similar to soil, microorganisms in the phyllosphere (both endo- and epiphytic) can be an important source of BVOCs. The phyllosphere contains bacteria, fungi, bryophytes, and lichens and are extremely diverse in tropical forests (Roy & Banerjee, 2018). Little is known about the relationship between the phyllosphere and BVOCs, but typical colonizers of the phyllosphere can emit and metabolise BVOCs (Farré-Armengol et al., 2016a; Kesselmeier et al., 1999; Rundell et al., 2015), therefore, the contribution of phyllospheric emissions to net ecosystem emission may be important given their abundance in rainforests (Roy & Banerjee, 2018). In addition to phyllospheric interactions, other processes at the foliar level can be a source
of BVOC emissions to the atmosphere, e.g. oxidation products formed interactions at the surface of leaves
(Jud et al., 2016) and adsorption/desorption onto/from surfaces.

Lastly, animals are a relatively unexplored source of BVOCs in Amazonian rainforests. In particular, arthropods such as ants and termites contribute 1/3 of the animal biomass in Amazonian upland forests, constituting several tonnes per hectare (Rinker & Lowman, 2004). Induced volatile emissions from animals, such as emissions from wood-eating termites of different social ranks (Himuro et al., 2011), or emissions of formic acid from ants for defensive purposes (Brütsch et al., 2017), have been reported. However, the types and magnitudes of the emissions from Amazonian ecosystems have not been really deepened since the reports by Zimmerman et al., (1982) or Khare et al., (1999).

202 The emission rates of BVOCs from all the aforementioned sources may vary considerably as a function of 203 driving factors. Temperature is a key driver of BVOC emissions, both for plants and soils, because it 204 promotes enzymatic reactions in pathways of BVOC biosynthesis and their diffusion in biological tissues 205 and soil substrates (Llusià & Peñuelas, 2000). For example, monoterpenes have been described as leaf "thermometers", increasing in emission with temperature (Jardine et al., 2017). Emissions of SQTs and 206 207 methanol are also temperature-dependent (Alves et al., 2016; Jardine et al., 2011c, 2016b) and isoprene has 208 been found to protect against thermal stress in Amazonia (Garcia et al., 2019). Soil temperature is also a 209 determinant because it increases decomposition and subsequent BVOC emission, but such processes have 210 not been adequately addressed.

211 Photosynthetically active radiation and its relationship to photosynthesis are essential drivers for 212 compounds emitted *de novo*, including isoprene and monoterpenes such as cis-β-ocimene (Alves et al., 213 2014; Harley et al., 2004; Jardine et al., 2013, 2015a, 2017; Kesselmeier & Staudt, 1999; Kuhn et al., 214 2002b, 2004a; Llusià & Peñuelas, 2000). For example, isoprene emissions from Hymenaea courbaril have 215 been strongly correlated with gross primary productivity under non-stress conditions (Kuhn et al., 2004b) 216 and protect plants under high solar radiation conditions (Garcia et al., 2019). Rapidly changing light 217 conditions, e.g. during a sudden thunderstorm, can also induce emissions of acetaldehyde and ethanol 218 (Holzinger et al., 2000). The effect of solar radiation on soils has not been well explored, but radiation 219 would both heat the surface and decrease the water content due to evaporation, though only about one third 220 of the radiation that arrives at the top of the canopy reaches the soil (Alves et al., 2016).

The availability of water within ecosystems affects BVOC production, whether as humidity, foliar uptake, or content in soil. Drought reduces stomatal conductance, decreases intracellular CO_2 mixing ratios and as a result, increases isoprene emissions (Pegoraro et al., 2004a), although this is dependent on the severity of drought (Niinemets, 2010; Otu-Larbi et al., 2020; Pegoraro et al., 2004b; Potosnak et al., 2014). GLV emission can be stimulated at high temperatures and under drought conditions (Jardine et al., 2015b;
Pfannerstill et al., 2018). The general reduction of BVOC emissions by plants due to drought depends on
both changes in plant physiology and to lower foliar biomass. Emissions from Amazonian soils depend on
moisture, because bacteria may become activated by water, with an emission burst after a large rainfall
event and an optimal pattern of emissions, where SQT emissions are the highest under moderate levels of
soil water (Bourtsoukidis et al., 2018). The soil water also affects the gas permeability of the soil. The soil
water content is therefore an essential parameter for BVOC production and release from Amazonian soils.

232 The atmospheric abundance of gaseous constituents is an additional regulator of BVOC emissions. The 233 mixing ratios of BVOCs in the atmosphere, particularly of oxygenated compounds that are bidirectionally exchanged (Niinemets et al., 2014), is driven by a compensation point (i.e. a compound is emitted if the 234 235 foliar concentration is higher than the ambient mixing ratios, otherwise the plant will take up the 236 compound), as has been observed in Amazonia (Jardine et al., 2011b; Kuhn et al., 2002a; Rottenberger et 237 al., 2004). The amount of oxidants in the atmosphere can induce emissions to quench oxidation (Jardine et al., 2011a, 2015a; Peñuelas & Llusià, 2001) and oxidise BVOCs generating oxidation products. Biotic and 238 239 abiotic stressors at the foliar level lead to the formation of reactive oxygen species (ROS), altering plant BVOC emissions coordinated to quench the ROS (Jardine & Jardine, 2016; Sampaio Filho et al., 2018). 240 241 Regional and global models currently have a module for deposition and another for emissions. For these 242 bidirectional exchange compounds such modules must be combined. One step forward in this matter is the 243 addition of a bidirectional exchange compound class in the MEGAN model (Guenther et al., 2012).

244 Increasing atmospheric CO₂ mixing ratios can affect BVOC emissions (Possell et al., 2005; Wilkinson et 245 al., 2009) directly, with higher biomass due to higher substrate availability (Lerdau et al., 1994), and thus 246 higher foliar surface area for BVOC emission, as demonstrated for monoterpene emissions (Staudt et al., 247 2001). However, no information on these processes is available for Amazonian ecosystems. These 248 increasing CO₂ mixing ratios can also indirectly suppress the emission of compounds such as isoprene 249 (Young et al., 2009), due to a competition of phosphoenolpyruvate (PEP) (Fasbender et al., 2018; Possell et 250 al., 2005; Rosenstiel et al., 2003), which at the same time may be offset by warming and increased growth 251 (Garcia et al., 2019; Sharkey & Monson, 2017).

Phenology, ontogeny, and canopy structure play important roles in BVOC emissions in Amazonian rainforests (Alves et al., 2018; Kuhn et al., 2004b; Wei et al., 2018). The quantity of BVOCs emitted by leaves varies with the stage of maturity as a consequence of resource availability (maturity) and defensive demand (bud break and early stages) (Kuhn et al., 2004b). For example, young *Vismia Guianensis* leaves emit higher quantity of methanol than mature leaves (Jardine et al., 2016b), whereas mature leaves *Vismia Guianensis* and *Eschweilera coriacea* have been found to higher emissions than other leaf ages (Alves et al., 2014, 2016; Jardine et al., 2016b); and light-dependent monoterpenes are emitted exclusively during the
period between bud-break and foliar maturity for *Hymenaea courbaril* (Kuhn et al., 2004b). Additionally,
canopy structure plays a role in BVOC dynamics as shown with high sesquiterpene emissions in the lower
canopy (Alves et al., 2016). Foliar flushing in Central Amazonia peaks during the transition from the wet to
the dry season, with a consequent peak in leaf litter and foliar turnover at the onset of the dry season
(Goulden et al., 2004; Huete et al., 2006; Roberts et al., 1998). This factor has been reported to be essential
for modelling BVOC emission in Amazonia (Alves et al., 2018).

Nutrient availability is another important driver of BVOC emissions (Fernández-Martínez et al., 2018). In this regard, the role of litterfall have recently been demonstrated to be important for supplying soil with nutrients that affect microbial composition in Amazonian ecosystems (Buscardo et al., 2018), but no information is available with respect to the effect of nutrient availability on BVOC dynamics at canopy level in Amazonia.

270 Biotic factors such as pathogens, herbivores, herbivorous predators, pollinators, and phyllospheric 271 organisms influence the emission rates of BVOCs. Plants emit BVOCs as defensive mechanisms or for signalling within plants and ecosystems (Šimpraga et al., 2019). Alterations to these biological controls can 272 273 thus affect the average BVOC emissions from plants. Little is known about the role of the phyllosphere in 274 BVOC dynamics in the Amazonian rainforest, but some possible implications can be derived from studies 275 elsewhere. The phyllosphere affects plant biogeography and ecosystem function, thereby affecting plant 276 physiology. The phyllosphere reduces the amount of photosynthesis by blocking light (Huete et al., 2006), 277 protects plants against heat (Stuntz et al., 2002), can retain water in the leaves (Coley et al., 1993; Stuntz et 278 al., 2002), fixes nitrogen, and protects plants against herbivores and abiotic stresses by the production of 279 phytohormones and other secondary metabolites (Roy & Banerjee, 2018). The phyllosphere is also 280 subjected to environmental conditions that can alter its role from beneficial to pathogenic. Such 281 environmental conditions can be leaching, changes in temperature, variations in exposure to sunlight, ROS 282 concentration (Bringel & Couée, 2015), the phyllospheric cover itself, effects of irrigation, cohabitation, 283 foliar texture, and phenology (Coley et al., 1993; Saikkonen et al., 1998).

Lastly, species diversity can influence the total emission of BVOCs, as has been reported for terpenoids, with substantial variation within species (Guenther, 2013). In Amazonia, an elevation gradient in isoprene emissions was attributed to the plant species distribution which differ topographically (Gu et al., 2017). The Amazonian rainforest contains a variety of ecosystems, from upland forests to flooded or seasonally flooded forests (see SI for more information). Differences in environmental conditions and adaptive strategies can be implied from differences in BVOC emission (Bracho-Nunez et al., 2012). Investigation on the effects of root anoxia on BVOC exchange is thus important for understanding the functioning of Amazonian ecosystems. Plants may react to flooding with fermentation, accompanied by emissions of ethanol and acetaldehyde to remove the large amounts of ethanol produced under root anoxia (Bracho-Nunez et al., 2012; Kreuzwieser et al., 1999; Rottenberger et al., 2008). The quantities and ratios of such emissions can indicate the metabolic and morphologic adaptations of flooded ecosystems (Parolin et al., 2004).

296 The vast amount of BVOC sources in the Amazonian rainforest (i.e. diverse species of insects, vegetation, 297 soil and phyllosphere microorganisms) is regulated by a plethora of biotic and abiotic emission drivers (i.e. 298 temperature, radiation, water and nutrient availability, ambient CO₂ and BVOC mixing ratios) leading to a 299 strong seasonality (Alves et al., 2016, 2018; Barkley et al., 2009; Kesselmeier et al., 2002b; Kuhn et al., 300 2004a; Yáñez-Serrano et al., 2015, 2018b). While increasing research addresses BVOC emissions over 301 single or combined emission drivers, the large-scale biodiversity and interplay of biotic and abiotic 302 processes remains challenging. Within such large ecosystems, ecological interactions may have an 303 important, but poorly explored, role in regulating the BVOC emissions of the Amazon rainforest.

Accepted

Table 1: Isoprene ambient mixing ratios reported in literature over Amazonian ecosystems in a within or above canopy and season basis. Data is organized by height with a line separating between within canopy mixing ratios and above canopy mixing ratios and between seasons. We considered 35m to be the average top of the Amazonian canopy. Values reported are in ppb and as presented in literature. They are usually means or medians. The values in parenthesis represent either standard deviation or interquartile range when available. Where only graphical data was available we used the software Plot Digitizer to derive means and standard deviations. The locations are classified according to the area where the measurements were taken, namely, Central Amazonia, Western Amazonia, South-western Amazonia, Guyana Shield and Central-east Amazonia. The techniques are PTR-MS: Proton Transfer Reaction Mass Spectrometer; GC-MS: Gas Chromatography Mass Spectrometer, GC-FID: Gas Chromatography Flame Ionization Detector and IEC: Ion-Exchange Chromatography. The seasons are specified together with the exact timing of sampling: Dry, dry season; Wet, wet season; D-t-W, transition from dry to wet season; and W-t-D, transition from wet to dry season. X stands for laboratory experiments or satellite retrievals.

For a more extended table with all reported values for Amazonian ecosystems in ppb, please refer to Table1 in S.I.2. Note that in some cases fluxes in mg m⁻² h⁻¹ are reported (these cases are identified by (flux), or by an *). The compounds are presented in the following order: Acetaldehyde, acetic acid, acetone, acetonitrile, acetylene, benzaldehyde, benzene, butanal, butane, butene, 1,4-cyclohexadiene, p-cymene, decanal, dichloromethane, diterpenes, dimethylsulphide, estragole, ethane, ethanol, ethylbenzene, ethylene, formaldehyde, formic acid, heptane, 1-heptene, 2-heptene, hexane, hexanol, hexene, hydrogen sulphide, hydroxyacetone (flux), isoprene (flux), linalool, methyl ethyl ketone (MEK), methanol, 2-methyl-2-butene, methyl chloroform, 2-methyl-furan, 3-methyl-furan, 6-methyl-5-heptene-2-one, methyl mecarptan, methyliodide, total monoterpenes, total monoterpene (flux), methyl vinyl ketone and methacrolein (MVK+MACR), MVK+MACR (flux), nonanal, octane, pentanal (flux), pentane, propane, propyne, total sesquiterpenes, total sesquiterpene (flux), styrene, sulphur dioxide, α -terpineol, toluene and m-, p- and o-xylene.

Isoprene

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Site	Season	Height	Height Isoprene	Technique Reference	Observations	
Sue		(m)	(ppb)		Nejerence	Coser viuons
Central Amazonia	Dry	0.05	3.13	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	0.5	3.77	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	1	2.77(0.4)	GC-FID	(Rasmussen & Khalil, 1988)	Daytime mean (11:00-15:00, LT). Canister sampling. Near ground level.

Western Amazonia	Dry	2	3.31	GC-MS	(Helmig et al., 1998)	Daytime mean.
Central Amazonia	Dry	4	5.35	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	0-30	2.40(1-5.24)	GC-FID	(Greenberg & Zimmerman, 1984)	Daytime mean. Canister sampling. Near ground to 30m
Central Amazonia	Dry	24	19.9(2.0)	PTR-MS	(Andreae et al., 2015)	Daytime mean.
Central Amazonia	Dry	24	7.62	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	30	2.65(1.39-3.38)	GC-FID	(Zimmerman et al., 1988)	24 h median. Sampling on teflon bag on tethered balloon (30m).
Central Amazonia	Dry	38	6.13	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	40	2.68(0.9)	PTR-MS	(Alves et al., 2016)	Daytime mean (10:00-14:00, LT).
Central Amazonia	Dry	40	7.94(1.67)	PTR-MS	(Wei et al., 2018)	Daytime mean.
Central Amazonia	Dry	45	5 max.	GC-MS	(Rinne et al., 2002)	Afternoon values. Fluxes 30 °C and 1000 µmol m ⁻² s ⁻¹ . Cartridge on Disjunct Eddy Accumulation.
Southwestern Amazonia	Dry	50	4.6(2.7)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Central Amazonia	Dry	51	3.4(1.8)	GC-FID	(Kuhn et al., 2007)	Daytime mean (10:00-15:00, LT). Cartridge on Relaxed Eddy Accumulation.
Central Amazonia	Dry	53	5.72	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	54	7.80	PTR-MS	(Rizzo et al., 2010)	Maximum at noon. Disjunct Eddy Covariance.
Central Amazonia	Dry	55	7.8(3.7)	PTR-MS	(Karl et al., 2007)	Daytime mean (12:00-14:00, LT). Disjunct Eddy Covariance.
Southwestern Amazonia	Dry	60	4.0(2.5)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Central Amazonia	Dry	79	4.81	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Dry	305	1.73(1.03-2.15)	GC-FID	(Zimmerman et al., 1988)	24 h median. Sampling on teflon bag on tethered balloon (305m).
Western Amazonia	Dry	91-1167	1.69	GC-MS	(Helmig et al., 1998)	Daytime mean.
Central Amazonia	Dry	100-1200	5.5(2.6)	PTR-MS	(Karl et al., 2007)	Daytime mean (10:00-11:30, LT). Mixed Layer Gradient.
Central Amazonia	Dry	150-1500	1.5(0.75)	GC-FID	(Rasmussen & Khalil, 1988)	Daytime. Canister sampling. Aircraft flights.
Central Amazonia	Dry	35-2000	2.27(0.38-4.08)	GC-FID	(Greenberg & Zimmerman, 1984)	Daytime mean. Canister sampling. Flights from treetop to 2 km.
Western Amazonia	Dry	1481-1554	0.16	GC-MS	(Helmig et al., 1998)	Daytime mean.
Central Amazonia	Dry	2000-17000	0.19(0.14-0.22)	GC-FID	(Greenberg & Zimmerman, 1984)	Daytime mean. Canister sampling. Flights from 2km to Tropopause.
Central Amazonia	Dry & D-t-W	2	1.54(1.37)	PTR-MS	(Jardine et al., 2011a)	
Central Amazonia	Dry & D-t-W	11	2.86(1.61)	PTR-MS	(Jardine et al., 2011a)	
Central Amazonia	Dry & D-t-W	17	2.86(1.53)	PTR-MS	(Jardine et al., 2011a)	
e e una i mazonia		24	2 51 (1 25)	DTP MS	(Jardine et al. 2011a)	

Southwestern Amazonia	Dry & D-t-W	08-52	9.5(4)	GC-FID	(Kesselmeier et al., 2002b)	Daytime mean (11:00-18:00, LT).
Central Amazonia	Dry & D-t-W	30	2.47(1.26)	PTR-MS	(Jardine et al., 2011a)	
Central Amazonia	Dry & D-t-W	40	2.36(1.26)	PTR-MS	(Jardine et al., 2011a)	
Central Amazonia	Dry & Wet	40	2.5(2.2)	PTR-MS	(Fuentes et al., 2016)	
Central Amazonia	D-t-W	40	2.65(1.33)	PTR-MS	(Alves et al., 2016)	Daytime mean (10:00-14:00, LT).
Central Amazonia	D-t-W	40	6.76(2.35)	PTR-MS	(Wei et al., 2018)	Daytime mean.
Southwestern Amazonia	D-t-W	50	3.4(1.2)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Southwestern Amazonia	D-t-W	60	3.0(0.5)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Central Amazonia	Wet	0	6.90	GC-MS	(Kesselmeier et al., 2000)	Daytime mean (15:00, LT).
Central Amazonia	Wet	0.05	0.57	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Wet	0.5	0.57	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Wet	1	3.2(0.9)	GC-MS	(da Silva, 2010)	Daytime mean (07:00-17:00, LT). Cartridge sampling.
Central Amazonia	Wet	1.5	4.33(2.02)	GC-MS	(Kesselmeier et al., 2000)	Daytime mean (09:30-16:00, LT). Outside forest.
Central Amazonia	Wet	1.5	3.01(0.85)	GC-MS	(Kesselmeier et al., 2000)	Daytime mean (09:30-16:00, LT). Inside canopy.
Central Amazonia	Wet	4	0.86	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central-East Amazonia	Wet	8.1	0.13(0.61)	GC-MS	(Jardine et al., 2016a)	Nighttime mean.
Central-East Amazonia	Wet	8.1	8.95(4.2)	GC-MS	(Jardine et al., 2016a)	Daytime mean.
Central Amazonia	Wet	10	4.6(0.94)	GC-MS	(da Silva, 2010)	Daytime mean (07:00-17:00, LT). Cartridge sampling.
Central Amazonia	Wet	20	6.17(1.03)	GC-MS	(da Silva, 2010)	Daytime mean (07:00-17:00, LT). Cartridge sampling.
Central Amazonia	Wet	24	2.29	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Wet	30	7.90	GC-MS	(Kesselmeier et al., 2000)	Daytime mean (14:00, LT).
Central Amazonia	Wet	38	2.39	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Wet	40	1.66(0.9)	PTR-MS	(Alves et al., 2016)	Daytime mean (10:00-14:00, LT).
Central Amazonia	Wet	40	4.66(2.21)	PTR-MS	(Wei et al., 2018)	Daytime medan.
Central Amazonia	Wet	50	6.3(1.32)	GC-MS	(Kesselmeier et al., 2000)	Daytime mean (12:00-16:00, LT).
Southwestern Amazonia	Wet	50	4.5(0.9)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Central Amazonia	Wet	53	1.85	PTR-MS	(Yáñez-Serrano et al., 2015)	
Southwestern Amazonia	Wet	60	4.0(1.2)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.

Central Amazonia	Wet	60	2.40	GC-MS/FID	(Batista et al., 2019)	Mean of overall dataset. Slope forest. Cartridge sampling with drones.
Central Amazonia	Wet	60	4.40	GC-MS/FID	(Batista et al., 2019)	Mean of overall dataset. Plateu forest. Cartridge sampling with drones.
Central Amazonia	Wet	79	1.69	PTR-MS	(Yáñez-Serrano et al., 2015)	
Central Amazonia	Wet	200-500	~3	GC-MS	(Kesselmeier et al., 2000)	24 hr mean. Sampling with thetered balloon.
Guyana Shield	Wet	400	~2.3	GC-FID	(Gregory et al., 1986)	Canister sampling.
Central Amazonia	Wet	500	~3	PTR-MS	(Martin et al., 2017)	During periods of clean conditions (above threshold).
Central Amazonia	Wet	200-1000	2.86(2.25-3.64)	GC-MS	(Greenberg et al., 2004)	Daytime median and interquartiles (12:00-15:00, LT). Cartridge sampling on tethered balloon.
Southwestern Amazonia	Wet	200-1000	6.89(2.78-7.73)	GC-MS	(Greenberg et al., 2004)	Daytime median and interquartiles (12:00-15:00, LT). Cartridge sampling on tethered balloon.
Central Amazonia	Wet	200-1000	0.74(0.6-1)	GC-MS	(Greenberg et al., 2004)	Daytime median and interquartiles (12:00-15:00, LT). Cartridge sampling on tethered balloon.
Guyana Shield	Wet	3000	3.30	PTR-MS	(Warneke et al., 2006)	
Guyana Shield	Wet	3000	1.70	PTR-MS	(Williams et al., 2001)	
Southwestern Amazonia	W-t-D	25	~4	GC-FID	(Kuhn et al., 2002b)	
Southwestern Amazonia	W-t-D	08-52	3.7(0.6)	GC-FID	(Kesselmeier et al., 2002b)	Daytime mean (11:00-18:00, LT).
Central Amazonia	W-t-D	40	4.17(1.81)	PTR-MS	(Wei et al., 2018)	Daytime mean.
Southwestern Amazonia	W-t-D	50	2.1(2.0)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Southwestern Amazonia	W-t-D	60	1.8(1.8)	GC-FID	(Aquino, 2006)	Daytime mean (11:00-16:00, LT). Canister sampling.
Central Amazonia	W-t-D	100	2.38(1.16)	GC-FID	(Kuhn et al., 2010)	Mean vertical profiles of isoprene and α -pinene from 8 background flights.
		Satellite	Х		(Barkley et al., 2009)	
		Satellite	Х		(Bauwens et al., 2016)	
Southwestern Amazonia	Dry	Leaf level	Х	GC-FID/MS	(Kuhn et al., 2004b)	
Southwestern Amazonia	Dry	Leaf level	Х	GC-FID/MS	(Kuhn et al., 2004a)	
		Leaf level	Х		(Bracho-Nunez et al., 2012)	
		Leaf level	Х		(Bracho-Nunez et al., 2013)	
		Leaf level	Х		(Garcia et al., 2019)	
		Leaf level	Х		(Harley et al., 2004)	
		Leaf level	Х		(Jardine et al., 2013)	

Leaf level Leaf level Leaf level Leaf level

Х

Х

Х

Х

(Jardine et al., 2016b) (Pegoraro et al., 2005) (Taylor et al., 2019) (Taylor et al., 2018)

4. Ecological interactions

BVOCs are assumed to play a strong role in ecological interactions in Amazonia (Figure 2). BVOC sources create volatile blends that can serve to identify the species of a recipient (Piva et al., 2019), allowing us to understand the nature of such communication (e.g. the identity and density of an attacking herbivore; Pinto-Zevallos et al., 2018), and can also direct the evolution of plant chemical diversity (Salazar et al., 2018).

Signalling within plants can be based on cascading signals of jasmonates, salicylates, and ethylene, but also
other types of BVOCs such as GLV, methanol or isoprenoids (Filella et al., 2006; Matsui, 2016; Matsui et
al., 2012; Seco et al., 2011). The function of their emission can be involved in: inducing the production of
BVOCs that can downregulate floral volatiles to produce BVOCs associated with fruit ripening, inducing
BVOC production for defensive purposes, or for individual plant coexistence (Kigathi et al., 2019;
Ninkovic et al., 2019). However, to our knowledge, these processes have not been investigated in
Amazonia.

Another type of signalling occurs between plants and other living organisms in ecosystems for 317 reproduction, defence, awareness of neighbouring plants, and modulating adaptation and evolution (Fine et 318 al., 2004; Laothawornkitkul et al., 2009; Salazar et al., 2018). Plants emit floral blends to attract specific 319 320 pollinators to aid reproduction. As rewards for pollination they offer pollen, and also resins, although the 321 latter more rarely, as for some Amazonian *Clusia* species (Bittrich & Amaral, 1997). Guarana trees emit 322 higher amounts of (E)- β -ocimene during night to attract nocturnal bees (Krug et al., 2018). The preferences 323 of bees for collecting pollen from other Amazonian plant species are nevertheless still unclear (Kaminski & 324 Absy, 2006). Furthermore, the role of phyllospheric floral bacteria has been pointed out to be of importance 325 for BVOC emissions (Peñuelas et al., 2014), although no information is available for the Amazonian 326 rainforest. Other types of communication associated with reproduction involve the production of fruit 327 odours for seed dispersal (Laothawornkitkul et al., 2009), for the seeds of a plant or of neighbouring plants 328 (Gfeller et al., 2019), particularly in Amazonia with seed-ant mutualism (Youngsteadt et al., 2009).

BVOCs emitted for defensive purposes have characteristic blends, in both abundance and chemical diversity, are species-specific (Courtois et al., 2009; Sharifi et al., 2018), serve an awareness role with neighbouring plants (De Moraes et al., 2001; Kessler & Baldwin, 2001), and can affect plant chemical diversity (Salazar et al., 2018). Such interactions can also occur between aboveground plants and roots, with important implications for BVOC interactions within the soil ecosystem (Delory et al., 2016). Herbivory in tropical rainforests has been estimated to consume 12-30% of the foliar area annually, with insects dominating foliar consumption, at 680 kg ha⁻¹ y⁻¹ (Rinker & Lowman, 2004). The role of BVOC emission in the Amazonian rainforest induced by herbivory is thus expected to be substantial but has notbeen characterized on regional scales.

338 Plants, including their roots, can emit BVOCs to attract predators of herbivores (Aartsma et al., 2017; 339 Sharifi et al., 2018). SQTs are good attractors of plant bodyguards (Rasmann et al., 2005) whereas isoprene 340 repels them (Loivamäki et al., 2008). For example, the Amazonian ant Allomerus octoarticulatus can be 341 attracted to damage from herbivory by its mutualistic plant Hirtella myrmecophila (Romero & Izzo, 2004), showing the complex interaction between plants and ants. In fact, the relationship between ants and plant 342 343 size in Amazonia can shape arthropod communities (Izzo & Vasconcelos, 2005). The extent of the damage 344 from herbivory feeds back on the defensive blends, as for cassava (Pinto-Zevallos et al., 2018), sometimes jeopardising the plant's protection (von Dahl et al., 2006). 345

Plants, microbes, and animals can emit BVOCs as direct toxins, deterrents, or repellents against herbivores
(De Moraes et al., 2001; Kessler & Baldwin, 2001; Laothawornkitkul et al., 2009; Prestwich, 1984;
Saikkonen et al., 1998; Sharifi et al., 2018). For example, Formicinae ants emit formic acid as a defensive
strategy (Brütsch et al., 2017) or GLV plant emissions can prime neighbouring plants against herbivores
(Engelberth et al., 2004). On the other hand, signalling can be negative, like isoprene that serves as a
negative cue, informing insects where not to forage (Gershenzon, 2008; Laothawornkitkul et al., 2008).

352 Protection against pathogens and sealing wounds are also drivers for BVOC emissions (Laothawornkitkul et al., 2009; Ninkovic et al., 2019; Šimpraga et al., 2019). For example, the biosynthesis of natural rubber 353 by Hevea brasiliensis is stimulated by wounding, releasing methyl jasmonate and ethylene (Duan et al., 354 355 2010), which is then oxidized during latex coagulation, sealing the wound (Piva et al., 2018). Nicotiana 356 tabacum plants emit methyl salicylate when pathogenic infections are present, inducing the expression of defensive genes in neighbouring plants (Shulaev et al., 1997) and phytopathogenes can turn plants into 357 358 sources of carbonyl sulphide (Bloem et al., 2012) rather than sinks as generally seen (Whelan et al., 2018). 359 The role of the phyllosphere in defence against pathogens has been established for Amazonia (Azevedo et 360 al., 2000; Gazis & Chaverri, 2015; Sia et al., 2013; Vaz et al., 2018), where pathogenic disease such as 361 anthracnose can be especially important for modulating the expansion and production of guarana 362 plantations (Bogas et al., 2015). BVOC emissions to protect against pathogens may thus be important for 363 identifying plant diseases.

Lastly, abiotic factors driving BVOC emission can also alter ecological interactions. Amazonian plants have been shown to protect themselves against excessive heat: by emitting isoprene (Jardine et al., 2016b) and distinct monoterpene species (Jardine et al., 2017), by oxidizing isoprene within leaf tissues and therefore releasing MVK+MACR (Jardine et al., 2011a), or by emitting GLV due to high temperatures and drought as part of induced senescence processes (Jardine et al., 2015b). There are other abiotic factors 369 altering ecological interactions. In Amazonia, rain can alter sesquiterpene speciation from soils 370 (Bourtsoukidis et al., 2018). Ozone can affect plant physiology (Loreto et al., 2004). BVOC oxidation can 371 disrupt plant olfactory cues (Vuorinen et al., 2004). Masking of BVOCs or ovipositional stimulants induced by herbivory may affect the range of the signal (Farré-Armengol et al., 2016b; Pinto et al., 2010). SQTs 372 373 emitted by soils and vegetation have been shown to quench ozone within the canopy of an Amazonian 374 rainforest (Bourtsoukidis et al., 2018; Jardine et al., 2011c). Thus, the abundance of ambient ozone regulates the distance over which the ecological BVOC signals are transported (Conchou et al., 2019). In 375 376 fact, given the toxicity of ozone, trees have been shown to emit reactive BVOCs to quench the ozone levels 377 (Bourtsoukidis et al., 2012). Additionally, ozone mixing ratios increase after biomass burning and urban pollution plumes in Amazonia (Wei et al., 2019), so these processes could be important for plant fitness 378 379 and ecological interactions in the future Amazonia. At last, atmospheric CO_2 increase is expected to alter ecological interactions either by increasing BVOC emissions due to higher substrate availability, decrease 380 381 BVOC emissions due to BVOC synthases competition in biosynthesis, or due to a modification of the 382 emitted compounds (Yuan et al., 2009).

While the role of BVOCs in ecological interactions is widely accepted, it is currently unknown how these will regulate the cumulative BVOC release from Amazonia. Disturbing established communication links among trees and between trees and insects may have unforeseen consequences for the ecosystem and species protection from pathogens. Anthropogenic activity alters ecological interactions; therefore, the rapidly changing atmospheric conditions may be a highly important piece of the puzzle in understanding the BVOC dynamics within Amazonia.

389 5. Atmospheric impact

390 The impact of Amazonian BVOCs on atmospheric chemistry is highly dynamic, with strong diel and 391 seasonal characteristics. The vast emissions of reactive BVOCs, in combination with high emissions of 392 other trace gases, insolation, high temperature and humidity, and high OH concentrations make the 393 Amazonian atmosphere a tropical photoreactor (Andreae, 2001). BVOCs emitted to the Amazonian 394 atmosphere are rapidly oxidized by OH radicals during the day and more slowly by ozone, or by NO_3 395 radicals at night (Paulson & Orlando, 1996). Some of the BVOCs emitted are very reactive, such as 396 isoprene towards OH (Nölscher et al., 2016) and SQTs towards O_3 (Jardine et al., 2011b). Isoprene, in 397 particular, can deplete tropical OH concentrations, thus indirectly affecting BVOC oxidation (Taraborrelli et al., 2012) and the lifetimes of climate relevant gases such as methane. 398

399 OH reactivity is a measurable parameter quantifying the total loss frequency of OH radicals in the 400 atmosphere, including BVOCs and other reactive trace gases. It can be used to assess how comprehensively 401 the reactivity of BVOCs can be accounted for in Amazonia. The difference between the calculated OH 402 reactivity vs. the observed OH reactivity indicates that there is a large portion of unexplained reactivity 403 (OH missing reactivity) possibly due to unmeasured compounds not accounted for in the calculation of OH 404 reactivity from the sum of the individual compounds. Missing reactivity measured at a single Amazonian rainforest location was strongly seasonal, being lower during the wet season (10 s⁻¹; missing OH reactivity 405 406 of 5-15%) and higher during the dry season (62 s⁻¹; missing OH reactivity of up to 79%), suggesting that 407 more unknown reactive compounds are present in the atmosphere during the dry season (Nölscher et al., 408 2016; Yang et al., 2016). Lower reactivity leads to a more oxidative atmosphere in the wet season, with 409 high deposition of oxidation products onto wet surfaces and possible enhanced microbial activity on foliar 410 surfaces that could influence the OH reactivity budget (Nölscher et al., 2016).

411 The gas-phase oxidation of BVOCs can form new particles by nucleation (Kirkby et al., 2016), or the 412 oxidised BVOCs can condense onto pre-existing particles, undergo heterogeneous reactions on particle 413 surfaces, be processed in clouds, or undergo further atmospheric degradation and deposition. 414 Photochemistry is an important Amazonian source of oxygenated VOCs (Yáñez-Serrano et al., 2015), 415 particularly for short-chain compounds (Seco et al., 2007). A recent study has demonstrated the strong role 416 of isoprene oxidation, which contributes ca. 20% of the organic aerosols (OAs) produced in Amazonia (Schulz et al., 2018). The contribution of other BVOCs to aerosol formation in the tropical rainforest is 417 more difficult to estimate, but SQTs have been estimated to contribute 0.4-5% to submicron OAs (Yee et 418 419 al., 2018). Shrivastava et al. (2019) recently reported that isoprene and terpenes can contribute almost half 420 of the Amazonian biogenic secondary organic aerosols (SOAs).

421 Aerosols have profound implications on the radiation balance locally, regionally and globally, either directly by scattering (Artaxo et al., 2013a; Kulmala et al., 2013) or absorbing solar radiation (Boucher et 422 423 al., 2013) or indirectly by impacting cloud and rain formation processes. Furthermore, they can boost 424 global primary production up to 25% for Amazonian ecosystems via diffuse radiation fertilization (Cirino 425 et al., 2014; Koren et al., 2012; Rap et al., 2018). Therefore, the presence of BVOCs influences the 426 associated albedo by modifying the number and size of cloud condensation nuclei (CCN) and cloud 427 droplets, affecting how the clouds reflect and absorb light (Boucher et al., 2013) and thus the radiation balance (Andreae & Crutzen, 1997; Artaxo et al., 2009; Sena et al., 2013) and hydrological cycle (Sheil, 428 2018). BVOCs and their role in SOAs, dominant type the fine-mode aerosol mass (Martin et al., 2010), 429 430 could ultimately affect the radiative cooling of the Earth (Ehn et al., 2014), particularly in Amazonia where 431 sulphur compounds are at low concentrations (Kirkby et al., 2016) and the radiation balance is driven by 432 organic aerosols (Artaxo et al., 2013a). A significant fraction of SOAs can act as CCN formed from 433 BVOCs in the Amazonian wet season (Pöschl et al., 2010). The low CCN particle concentrations lead to larger cloud droplets producing clouds at lower altitudes that have higher rain efficiency (Rosenfeld, 1999), 434 435 intensifying the role of BVOCs in aerosol and cloud dynamics. The formation of new particles at the planetary boundary layer has not been observed in Amazonia. BVOC species are instead elevated by deep
convection and converted into condensable species in the upper troposphere, forming new aerosol particles
and increasing the loading of OAs, which can then be transported down to the planetary boundary layer
(Andreae et al., 2018; Schulz et al., 2018; Figure 2).

440 BVOCs can also affect radiative forcing by their degradation in the atmosphere (with CO₂ as an end product) and by their perturbation of the lifetime of other greenhouse gases such as methane, ozone 441 442 (Guenther, 2002), and further reactive gases such as CO (Collins et al., 2002). This is estimated to add 0.16 Pg C y⁻¹ to the amount of global atmospheric CO₂ (Suntharalingam et al., 2005), which can increase CO₂ 443 444 levels by 0.07 ppm y⁻¹, and can have a significant impact on the results of inverse model investigations of 445 CO₂ sources. Because BVOCs deplete OH faster than methane (for which the primary sink is OH), methane 446 will have a longer lifetime in the atmosphere (Collins et al., 2002). Alternatively, tropospheric ozone is 447 formed via BVOC oxidation when sufficient NOx is available, adding to the radiative forcing of 448 greenhouse gases.

449 NOx levels are low (ca. 0.4 ppb) in the atmosphere over the undisturbed Amazonian rainforest (Liu et al., 2016), particularly during the wet season when air masses arrive predominantly from the northeast over 450 451 mostly pristine rainforest (Pöhlker et al., 2019). NO from soils that reacts with O_3 to form NO₂ inside the canopy is the main source of NOx. This NO₂ is deposited on the leaves of the very dense rainforest canopy, 452 453 reducing the contribution of soil NO emissions to the levels of NOx above the canopy (Jacob & Wofsy, 454 1990). The direct loss by reaction of O₃ with BVOCs outcompetes O₃ formation under these pristine 455 conditions, thereby leading to a decrease in O₃ (Jacob & Wofsy, 1990; Scott et al., 2018). O₃ budgets are 456 thus mainly controlled by transport from above (downdrafts from convective storms (Dias-Júnior et al., 457 2017; Gerken et al., 2016) and deposition to the canopy, with little net effect from photochemistry (Jacob & Wofsy, 1990). 458

459 NOx levels increase significantly during the dry season due to changes in wind direction from the easterly 460 and southeasterly fetch regions, transporting considerable amounts of pollution from burning biomass and 461 human activities (Bela et al., 2015; Pacifico et al., 2015). Consequently, O₃ levels also increase during this 462 season (8-15 ppb in the wet season, 15-50 ppb in the dry season at midday) (Andreae et al., 2015; Artaxo et 463 al., 2013a, Figure 5). The concentrations of trace gases and particles are thus strongly seasonal. In fact, an 464 increase in NOx mixing ratios (to >1 ppb) can affect the oxidative pathways (i.e. HO₂ or NO pathway; Liu et al., 2016) of BVOC peroxy radicals, thus altering photochemistry, even in this region where NOx levels 465 466 are generally low. The central Amazonia plume has been reported to significantly increase O_3 levels by 35% at 70 km downwind of Manaus, whereas NOx levels increased by <20% relative to remote areas, 467 indicating the strong role of BVOC oxidation in forming O₃, in areas where NOx levels have increased due 468

to pollution (maxima of up to 6 ppb; Wei et al., 2019). Enhanced urban pollution and plumes from burning
biomass alter BVOC emissions by inducing further emissions of highly reactive BVOCs (Bolsoni et al.,
2018) and thus, altering the dynamics of natural aerosols, with an increase in SOAs formed by BVOC
oxidation (Shrivastava et al., 2019). The burning of biomass can profoundly alter SOA composition,
although increases in organic submicron particulate matter during the dry season are also due to increased
BVOC emissions combined with reduced deposition (de Sá et al., 2019; Williams et al., 2016).

- Canopy structure is another very important parameter in determining the impact of BVOCs in the 475 476 atmosphere. The dense and tall canopy of the Amazonian rainforest can stratify air masses, with an 477 intermittent coupling of the upper canopy and the atmosphere above. Air is poorly mixed within the 478 canopy, particularly at night when conditions are stable, with occasional exceptions of periods with bursts 479 of intermittent turbulence (Freire et al., 2017). The lower part of the canopy is characterized by a thermal 480 inversion, hindering mixing between the canopy and the forest floor (Nölscher et al., 2016; Santana et al., 481 2018). This feature can affect the residence times of emitted BVOCs and their turbulent transport out of the 482 forest canopy (Bakwin et al., 1990; Fitzjarrald et al., 1988; Gerken et al., 2017). BVOCs can thus remain 483 trapped within the canopy, becoming oxidised before reaching the atmosphere above. For example, 50-70%of O_3 reactivity within the canopy has been attributed to the ozonolysis of highly reactive SQTs 484 485 (Bourtsoukidis et al., 2018; Jardine et al., 2011c; Yee et al., 2018).
- 486 Isoprene oxidation by OH produces, among many other products, isoprene peroxide radicals (ISOPOO). 487 These radicals can react with HO₂ to form isoprene hydroxyhydroperoxides (ISOPOOH) or with nitric 488 oxide (NO) to produce mostly methyl vinyl ketone and methacrolein. Both pathways (HO₂ and NO) are 489 important in Amazonia, despite their strong seasonality (Liu et al., 2016). The formation of isoprene 490 epoxydiols can lead to the growth/formation of SOAs, mostly in the upper troposphere (Schulz et al., 491 2018). ISOPOOH levels can nevertheless decrease in polluted conditions (Liu et al., 2016), with 492 implications for OA levels, because higher NOx levels can suppress the production of SOAs derived from 493 isoprene epoxydiols (IEPOX) (de Sá et al., 2017; Schulz et al., 2018). Isoprene was previously thought to 494 yield very low quantities of condensable material under low NOx levels (Pandis et al., 1991). More 495 recently, gas-phase IEPOX, second-generation products of isoprene oxidation under RO₂+HO₂ chemistry 496 (Surratt et al., 2010), and reactive uptake have been proposed as an additional pathway. The IEPOX may be 497 responsible for the production of observed SOA isoprene tracers, such as 2-methyltetrols found in the 498 particle phase of Amazonian samples (Carlton et al., 2009; Claeys et al., 2004; Kroll et al., 2005).

In summary, BVOCs can have very strong impacts on the atmospheric chemistry over Amazonia. BVOC emissions determine oxidant regimes, as they modulate the major atmospheric oxidant, OH, and they can lead to pollutant production (e.g. tropospheric ozone) when the pristine Amazonian conditions are altered.

502 6. Depositional processes

503 Deposition of BVOCs can be divided into wet and dry deposition onto surfaces, biological uptake, and 504 physicochemical processes at the leaf-atmosphere interface (Figure 2). In Amazonia, wet deposition can be 505 an important sink for trace gases. Polar compounds are particularly well washed off by rain. Moreover, 506 BVOC oxidation products can be formed in the water layer of leaves and other surfaces. This process is 507 especially important during the wet season and periods of rain and contributes to the observed general decrease in the concentrations of trace gases. For example, organic acids such as formic and acetic acid can 508 509 contribute up to 90% of rain acidity in Amazonia (Andreae & Andreae, 1988; Jardine et al., 2011b). Dry 510 deposition depends on the movement of air, the vertical stability of the boundary layer, the chemical 511 properties of the species, and the nature of the depositional surface (Stickler et al., 2007), which may be 512 important in Amazonia, particularly at night with 78% of isoprene lost to surface deposition (Wei et al., 513 2018). Further information is needed to understand dry deposition in Amazonian ecosystems, which may 514 have been underestimated as has been demonstrated over a dense tropical rainforest in Costa Rica (Karl et 515 al., 2004).

516 Vegetation takes up compounds via stomatal or non-stomatal processes, where re-metabolisation can occur (Matsui, 2016). The ambient mixing ratio of a compound regulates its uptake or emission depending on the 517 compensation point. Compensation points for Amazonian ecosystems have been reported to be <0.6 ppb 518 519 for acetaldehyde and formaldehyde (Rottenberger et al., 2004) and <0.3 for formic and acetic acid (Kuhn et 520 al., 2002a) based on leaf level measurements. Data estimated on an ecosystem level were found to range 521 around 1.4 and 2.1 ppb for formic and acetic acid, respectively (Jardine et al., 2011b). In view of the 522 possible bio-facilitation for plant VOC uptake (Matsui, 2016), more information is needed to understand 523 the function of bi-directional BVOC exchange by Amazonian ecosystems.

524 Bacteria are another sink or source for BVOCs. They can take up BVOCs as a source of carbon (Ginkel et 525 al., 1987) when levels are not toxic and can detoxify BVOCs above toxic levels (Marmulla & Harder, 526 2014). For example, isoprene can be taken up by tropical soils (Cleveland & Yavitt, 1997), depending on 527 the atmospheric mixing ratio of isoprene and with decreasing rates of uptake under drought conditions 528 (Pegoraro et al., 2005). Bacterial communities in the phyllosphere may also be a BVOC sink (Farré-Armengol et al., 2016a). Phenol degradation has been observed in phyllospheric communities elsewhere 529 530 (Sandhu et al., 2007). All these phyllospheric sink processes for BVOCs could be especially important in 531 Amazonia, given its abundant phyllospheric communities.

Processes taking place at the leaf surface such as oxidation and adsorption/desorption are important for ecological interactions, plant physiology, and accounting for gross and net BVOC emissions. Oxidation can occur in the stomatal cavity or at the leaf surface as it serves to quench ROS damage in vegetation. Examples have been reported for monoterpenes (Loreto et al., 2004), by semi-volatile organic compounds
exuded by glandular trichomes (Jud et al., 2016), and specifically in Amazonia for isoprene (Jardine et al.,
2011a). This oxidation can lead to the production of oxygenated BVOCs such as ketones and aldehydes
(Jardine et al., 2011a; Jud et al., 2016; Pinto et al., 2010). This within-leaf oxidation process is of particular
importance for dry deposition estimates worldwide. When fast metabolic conversion of oxygenated BVOCs
was incorporated into a global chemistry transport model, Amazonia showed an increased annual dry
deposition flux of 75% (Karl et al., 2010).

Adsorption/desorption can occur on surfaces, such as plant cuticles (Müller & Riederer, 2005) and water films and may be important in humid places such as Amazonia, even though their functioning is not yet fully understood. Water layers on the upper surfaces of leaves may strongly influence the absorption of water by leaves, stomatal conductance, pathogenic infection, nutrient leaching, and adsorption of pollutants (Aryal & Neuner, 2010). Oxygenated BVOCs may thus be adsorbed onto water films and are then released upon the evaporation of the films. This process may occur in wet environments such as Amazonia.

548 In summary, BVOC deposition is dependent on many factors, including the nature and concentration of the compound and the existent meteorology. Despite high uncertainties, in an attempt to quantify the 549 550 magnitude of BVOC deposition, we used approximations from literature. Dry deposition has been 551 estimated to occur onto leaf surfaces as well as to soil surfaces with shares of 70% and 30%, respectively 552 (Karl et al., 2010). These estimates are expected to be higher for leaf surfaces in Amazonia due its larger 553 leaf area. On the other hand, in Amazonia, rain wet deposition occurs 90% to soil surfaces, with only 10% 554 being intercepted by vegetation (Czikowsky & Fitzjarrald, 2009). In addition, fog wet deposition is 555 expected to occur at high percentage values (Shimadera et al., 2011), although no estimates have been 556 found so far for Amazonia. A proper characterization of Amazonian BVOC deposition flux could result in 557 higher estimates of gross BVOC production, larger than previously thought. The aforementioned processes 558 may account for active and partly unexplored sinks, which have to be considered in regional and global modelling exercises. 559

560

7. Climate change, land-use change, and feedbacks

Amazonian ecosystems may be moving towards tipping points. For example, a temperature increase of 4°C or deforestation exceeding 40% of the forested area (Nobre et al., 2016) due to human intervention as landuse change possibly lead to savannisation (Marengo et al., 2011) and limit rainfall in synergistic interactions with the impacts of climate change.

565 Climate change is expected to alter Amazonian ecosystems, since increased warming will be combined
566 with decreased precipitation (IPCC, 2013). Accompanying effects are drought associated with El Niño–
567 Southern Oscillation (ENSO) anomalies and changes in the surface temperature of the Atlantic Ocean

(Leitold et al., 2018), increased frequency and intensity of heat waves (Perkins-Kirkpatrick & Gibson,
2017), and increased duration and intensity of dry (Adams et al., 2017) and intensity of wet (Leite-Filho et
al., 2019; Mori and Becker, 1991) seasons. Increasing temperatures and drought (Aragão et al., 2018;
Feldpausch et al., 2016; Leitold et al., 2018; Li et al., 2008; Marengo et al., 2016; Pfannerstill et al., 2018;
Yang et al., 2018) and intensified dry seasons (Sena et al., 2018), which are slowly drying Amazonia
(Esquivel-Muelbert et al., 2018), have already been observed.

These conditions of thermal and drought stress have multiple effects in Amazonian ecosystems. An 574 575 intensified dry season can particularly increase the frequency of fires (Silva Junior et al., 2019), reduce net 576 primary productivity, and accelerate canopy turnover (Leitold et al., 2018). Drier conditions can lead to an overall decrease in BVOC production (although drought may induce transient GLV emission (Jardine et al., 577 578 2015b) (Figure 6). Other effects are higher irradiance due to lower cloud cover (Sena et al., 2018) and 579 higher temperatures, which ultimately lead to a net increase in BVOC production (Jardine et al., 2017; 580 Pfannerstill et al., 2018) of both BVOCs associated with stress (Jardine et al., 2015b) and other unknown 581 highly reactive compounds (Pfannerstill et al., 2018). If threshold limits for heat and radiation are 582 exceeded, increased tree mortality will naturally reduce BVOC emissions. Altered ecological interactions (e.g. insect outbreaks or altered floral compositions) and increasing pollution can modify BVOC emissions, 583 584 with unforeseen consequences. However, an intensified wet season, could lead to stronger storms (Nelson 585 et al., 1994) and floods (Barichivich et al., 2018), increasing BVOC production associated with root anoxia. Tree mortality associated with storms (Aleixo et al., 2019) have potential implications for atmospheric 586 587 chemistry because of increased BVOC emission associated with anoxia from floods.

These effects ultimately lead to increased tree mortality (Aleixo et al., 2019; Brando et al., 2019) aggravated by deforestation, and, therefore, to decreased BVOC production due to biomass reduction, which further feeds back on a changing climate by altering ecological interactions and aerosol dynamics. Additionally, the loss of biodiversity (Nobre et al., 2016) caused by climate change (Chapin III et al., 2000) and deforestation (Barlow et al., 2007) can alter ecological interactions and associated BVOC emissions.

593 Agricultural practices, including but not limited to biomass burning, and urbanisation are the two main 594 impacts of land-use change affecting Amazonia. Twenty percent of the original forest in the Brazilian 595 Amazonia has been cleared in the last three decades (Bullock et al., 2020; Davidson et al., 2012), 596 particularly from the so-called arc of deforestation in southern and southwestern Amazonia (Leite-Filho et 597 al., 2019; Pöhlker et al., 2019), with a sharply increasing trend during the past several years. Deforestation 598 has implied substantial environmental changes (Davidson et al., 2012) and forest disturbance (Bullock et 599 al., 2020), such as water stress due to shallower roots (Huete et al., 2006) that causes an energy imbalance, 600 loss of biodiversity, changes to carbon storage (Huete et al., 2006), and an overall decrease in BVOC

601 emissions due to the loss of biomass (Scott et al., 2018). Biomass burning also profoundly affects the 602 oxidation pathways of BVOCs and increases emissions of air pollutants (Bulbovas et al., 2014) by clearing 603 land mainly for soy production (Bulbovas et al., 2007), oil palm production (Jardine et al., 2016a) and 604 cattle ranching (Figure 5). Soy crops are very sensitive to tropospheric ozone (Bulbovas et al., 2007), so 605 higher ozone mixing ratios will feed back to more deforestation to counteract decreased soy productivity 606 (Pimentel, 2011).

Increasing urbanisation (Richards & VanWey, 2015) and rural development (Nobre et al., 2016) are also 607 608 drivers of land-use change in Amazonia. The urban population of the region has increased from about 1 609 million to 11 million in the last 50 years (1960–2010) (Medeiros et al., 2017). In central Amazonia, the city of Manaus has 2 million inhabitants, and pollution sources include vehicular emissions, refinery operations, 610 611 and electricity generation. Even though anthropogenic emissions are lower than for other regions around 612 the world, Amazonian air chemistry is highly sensitive, amplified by the high solar irradiance and 613 concentrations of water vapour in an environment with plentiful BVOC emissions. NOx and ozone 614 production are much higher in urban plumes (Kuhn et al., 2010; Trebs et al., 2012; Wei et al., 2019) and 615 have profound impacts on atmospheric chemistry and physics by increasing biogenic SOA production by 60-200% (Shrivastava et al., 2019) and altering gaseous (Liu et al., 2016) and particle (de Sá et al., 2017) 616 chemical pathways. A programmed change from fuel oil and diesel to natural gas is nevertheless projected 617 618 to improve urban air quality (Medeiros et al., 2017).

619 At a global scale BVOC emissions may increase in a warmer climate (Peñuelas & Staudt, 2010) due to 620 higher plant metabolism and diffusion, but in Amazonia, the effects of climate could increase the emission 621 of stress-related BVOCs and the effects of land-use change could lead to a decrease in BVOCs due to loss 622 in biomass. An increase of BVOCs due to warmer temperatures would lead to more SOA having a cooling 623 impact through the direct light scattering or indirectly by the increase in cloud cover. This would lead to a 624 negative radiative forcing. Additionally, more SOA would lead to more diffuse radiation, boosting GPP 625 (Rap et al., 2018) and thus the uptake of atmospheric CO₂. On the other hand, an increase in BVOCs would 626 lead to an increase of the CH₄ lifetime. In the presence of sufficient NOx, an increase in tropospheric ozone 627 could be a consequence resulting in a warming effect. A decrease in BVOC emissions, however, would 628 have opposite effects on the radiative balance. Moreover, the net effects of altered ecological interactions 629 (Niederbacher et al., 2015), higher oxidation capacity, changes in species composition, and increased flooding on BVOC responses remain unknown for Amazonian ecosystems. Unfortunately, there is almost 630 631 no information about the impact of BVOCs in the Amazonian hydrological cycle, with effects for the rest 632 of the South American continent (Boers et al., 2017; Rocha et al., 2018). Thus, a substantial change in Amazonian BVOC emission may have strong climatic impacts at a regional and even at global levels, but 633 634 there are large uncertainties in current estimates.

635 8. Recent evolution of isoprene emissions in Amazonia

636 We investigated the evolution of isoprene fluxes in Amazonia using modelled isoprene emissions from the CAMS-GLOB-BIO dataset (Granier et al., 2019), and satellite-based estimates (Bauwens et al., 2016) for 637 638 2005-2014. The CAMS-GLOB-BIO global dataset is based on the Model of Emissions of Gases and Aerosols from Nature (MEGANv2.10; Guenther et al., 2012) and includes monthly isoprene emissions at 639 640 0.5°x0.5° resolution. The MEGAN model was driven by re-analysed ERA-Interim meteorological fields (Dee et al., 2011). Global annual land cover was provided by the ESA-CCI dataset (Climate Change 641 642 Initiative of the European Space Agency, ESA, 2017), processed in order to be consistent with plant 643 functional types (PFT) used in the MEGAN model (see S.I.3). On the other hand, for the satellite data we 644 used formaldehyde columns from the OMI satellite sensor and an inverse modelling framework (see S.I.4.). 645 Both techniques were used successfully for estimating isoprene fluxes in this region in the past (Barkley et 646 al., 2009; Pacifico et al., 2011). Seasonal and inter-annual variations of the modelled and satellite-based 647 isoprene fluxes over the Amazonian region (5°N-13°S, 46-77°W) are shown in Figure 7. Both approaches 648 display similar magnitudes and seasonal patterns, with slightly higher fluxes for modelled isoprene. The 649 decrease of isoprene fluxes during the wet season is observed in the complete time series of modelled isoprene (1992-2015) (Figure S.I.2). Since biomass burning during the dry season affects the formaldehyde 650 columns, only wet season data are used for the trend calculation, but it has to be noted that both modelled 651 652 and satellite isoprene fluxes show a decrease for the reported period. The decrease of isoprene fluxes during 653 the wet season can be partly attributed to large and rapid changes in land cover (Acosta Navarro et al., 2014). Even though the rate of deforestation decreased in the period of 2005-2014 (Artaxo et al., 2013b), 654 655 studies point out that drought-related fires may have increased during this period (Aragão et al., 2018). In fact, satellite observations suggest a widespread forest degradation due to deforestation (Bullock et al., 656 657 2020).

The modelled isoprene flux indicates an increase of isoprene emissions during the dry season for 1992-2015, with peaking fluxes in El Niño years (1997, 2015; Figure S.I.2). Given that sesquiterpenes are temperature-dependent and emitted as a plant response to environmental stresses (Zhao et al., 2017), we have calculated the sesquiterpene to isoprene ratio as a proxy of environmental stress (Figure 8). The ratio significantly increases over time, suggesting that Amazonian forest is undergoing increased heat stress due to climate warming.

These results underline the combined effect of climate change and land use change affecting BVOC fluxes.
During the dry season, strong environmental stresses to Amazonian vegetation occur at higher intensity and
frequency over time. This view is supported by the increasing ratio of sesquiterpene to isoprene fluxes.

667 During the wet season, the decrease in isoprene emissions may be partly explained by the rainforest loss668 due to deforestation, although further analysis is needed to prove this hypothesis.

669

9. Research priorities

670 Kesselmeier et al. (2009) identified several research priorities that need to be addressed to better understand 671 BVOC dynamics in Amazonia. However, some of them remain unaddressed even one decade later. 672 Intensive efforts towards integrated investigations of emissions, chemistry, transport, and cloud processes, 673 with campaigns such as Brazilian-Air 2010, GOAMAZON (2014-2015), ACRIDICON (2014), and ATTO 674 (2012-present), which comprised observations from tower-based measurements (at TT34, K34, ATTO) and 675 aircraft-based studies (ACRIDICON and GOAMAZON) have been made or planned for the near future 676 (CAFÉ-Brazil 2020). These studies provided essential information, particularly in understanding the fate of 677 emitted isoprene in the atmosphere. Many studies have now addressed a wide range of biogenic VOCs, 678 including oxygenated BVOCs, aromatic compounds, sulfurous compounds, and oxidation products (Alves 679 et al., 2016; Bourtsoukidis et al., 2018; Jardine et al., 2011b, 2011a, 2014, 2015a, 2015b, 2017; Liu et al., 680 2016; Paralovo et al., 2015; Yáñez-Serrano et al., 2015, 2016, 2018b; Yee et al., 2018), although new studies should aim to characterize unknown reactive compounds to better understand atmospheric 681 682 reactivity, particularly in the dry season (Nölscher et al., 2016; Pfannerstill et al., 2018). Progress has been 683 made in understanding of the impact of BVOCs on oxidant and aerosol distributions (Andreae et al., 2018; 684 de Sá et al., 2017, 2019; Liu et al., 2016; Nölscher et al., 2016; Pfannerstill et al., 2018; Schulz et al., 2018; 685 Shrivastava et al., 2019), the processes controlling the seasonal (Alves et al., 2016, 2018; Wei et al., 2018; 686 Yáñez-Serrano et al., 2015, 2018a) and spatial (Barkley et al., 2009, 2011; Bauwens et al., 2016; Salazar et 687 al., 2018; Taylor et al., 2018, 2019) variations of BVOCs, flooding (Bracho-Nunez et al., 2012), and 688 ecological interactions (Salazar et al., 2018). However, more process-based studies are essential, such as 689 those using laboratory and enclosure methods, to understand mechanistically what is happening at foliar 690 surfaces, including phyllospheric processes, BVOC sink processes, relationships between BVOCs and the 691 carbon and water cycles, ecological interactions, and the response of BVOCs to climate change and 692 deforestation in Amazonia. Model simulations could thus become more process-based, reducing the 693 uncertainty of predictions. For example, strong efforts are currently being made to understand BVOC 694 variation across plant populations beyond plant functional type, which is too broad to represent variability 695 in Amazonian ecosystems (Taylor et al., 2018).

696 Understanding how BVOC emissions would affect CCN and rain formation, including rain acidity (Jardine
697 et al., 2011b), is important as clouds can have an effect on the radiative forcing by reflecting radiation and
698 by increasing diffuse radiation thereby increasing GPP, as well as having impacts on the hydrological cycle
699 (Koren et al., 2012). But the effect of increased flooding, as a response to climate change, must also be

700 understood at an ecosystem level, particularly the role of anoxia BVOCs in CH₄ and CO₂ cycles in flooded 701 ecosystems. More effort is needed to obtain a more regional and long-term vision of BVOCs in Amazonia, 702 which can only be approached with synergistic investigations of different Amazonian ecosystems, to understand the timing and duration of the response dependency of BVOC dynamics. This review 703 704 demonstrates the necessity of understanding the signals of BVOC feedback loops in response to climate 705 and land-use changes in Amazonia, for which understanding the potential impacts of the role of BVOCs in 706 ecological interactions is crucial, a subject rarely studied in Amazonia despite its enormous biological 707 productivity and biodiversity. Understanding how these interactions can be modulated by climate change is 708 also critical. Some ecological theories have been proposed to account for hyperdominant species in 709 Amazonia (ter Steege et al., 2013), which study the evolution and distribution of defensive compounds as a 710 function of species composition. For example, *Protium* genus produces large quantities of phenolics, which may increase colonisation of its species (Salazar et al., 2018). This approach may be used to study the 711 712 ecological interactions of BVOCs in Amazonia. Climate change and pollution increase the oxidative 713 capacity of the Amazonian atmosphere (Nölscher et al., 2016; Pfannerstill et al., 2018; Yáñez-Serrano et 714 al., 2015), but the signals of the BVOC responses in the feedback loops remain unknown. Similarly, the 715 signal of the BVOC response to an intensified wet season remains unknown. The responses of BVOCs to global change in Amazonia, either as a direct response to stress (e.g. increase drought, increase in UVB or 716 717 nutrient availability) or as an indirect response due to the extinction of BVOC sources (e.g. tree mortality or 718 agricultural expansion), are key research priorities for understanding how Amazonian ecosystems function and how they will be modified in the future. This review suggests that warming and associated drying, 719 720 combined with the rate of deforestation, deforestation and associated drying is and will be the most determinant in the fate of BVOCs in Amazonia (both for chemistry and physics), impacting the radiative 721 forcing of the atmosphere and the hydrological cycle. Deforestation is a major threat to Amazonia, with 722 723 >10000 km² between the period between August 2018 and July 2019, the highest annual loss since 2008 724 (Barlow et al., 2020). Even the most optimistic climatic scenarios where forest protection policies are 725 implemented predict substantial deforestation in Amazonia (Pöhlker et al., 2019; Soares-Filho et al., 2006). This trend indicates a major loss of forest biomass, potentially reaching threshold levels with difficult 726 727 recovery. This loss will affect not only the tropical and extra tropical carbon and water cycles, but also the 728 BVOC exchange between the surface and the atmosphere. Actions such as large-scale forest restoration 729 (Bastin et al., 2019) and the promotion of services (Šimpraga et al., 2019) provided by Amazonian 730 ecosystems may therefore help to mitigate these impacts of deforestation. Human action will thus delineate 731 also the future of BVOCs in Amazonia.

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11. Additional Information

753 Competing Interests: The authors declare no competing interests.

754 Data accessibility: Data is available upon request to corresponding author.

755 **12. References**

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1784	Research 93(D2) 1407 https://doi.org/10.1029/ID093iD02p01407
1785	
1786	13. Figures



Figure 1: Map of Amazonia with the distribution and frequency of published BVOC measurements. The centres of the
circles represent the locations of the measurements, and the sizes of the circles represent the number of published
studies of BVOCs. This graph was derived from the information of Table 1 and S.I.1.



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Figure 2: Schematic showing the BVOC dynamics in a pristine Amazonian atmosphere, including sources and
interactions, BVOC oxidation in the atmosphere, and subsequent aerosol production influencing cloud droplet
formation and cloud dynamics. CCN stands for cloud condensation nuclei and SOA stands for secondary organic
aerosol.



Figure 3: Whiskers box plot of all measured ambient concentrations (in ppb) of BVOCs in Amazonia to date. The
ends of the box are the upper and lower quartiles showing the interquartile range, and the median is marked by a
horizontal line inside the box. The whiskers are the two lines outside the box that extend to the highest and lowest
observations.



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Figure 4: Whiskers box plot of all measured ecosystem fluxes (in mg m⁻² h⁻¹) of BVOCs in Amazonia to date. The
ends of the box are the upper and lower quartiles showing the interquartile range, and the median is marked
by a horizontal line inside the box. The whiskers are the two lines outside the box that extend to the highest
and lowest observations.

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000 cm⁻³ Aerosol Condensable vapors particle CCN SOA (New particle formation) Downdrafts SOA (Particle growth) Condensable vapors Low NOx High NOx $H_2O+NO \rightarrow NO+O \rightarrow O+O_2+M \rightarrow O_3+M$ BVOC+OH $\rightarrow RO_2 + NO \rightarrow RO$ $HO_2 + O_3 \rightarrow OH + 2O_2$ BVOC+OH $\rightarrow RO_2 \rightarrow ROOH$ Gas-p oxidation CO₂ CO₂ Altered BVOC + AVOC

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Figure 5: Schematic showing BVOC dynamics in a polluted Amazonian atmosphere, including BVOC sources and interactions, interaction with anthropogenic volatile organic compounds (AVOCs), BVOC oxidation in the atmosphere, and subsequent aerosol and cloud dynamics. CCN stands for cloud condensation nuclei and SOA stands for secondary organic aerosol.

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1815 Figure 6: Proposed feedback loops for the impacts of climate and land-use changes in Amazonian BVOCs and how 1816 these impacts will feed back into the system. The dashed arrows represent the situation where threshold limits are 1817 reached (i.e. heat can stimulate BVOC emissions until the point where heat is detrimental to plant fitness, thereby

1818 decreasing BVOC emissions). Red colour stands for promoting and blue colour for inhibiting. Grey arrows indicate
1819 inconclusive/unknown effects. LULCC stands for Land Use Land Cover Change.



Figure 7: Monthly top-down isoprene flux estimates based on global model simulations using MEGAN (top) and
based on the OMI satellite formaldehyde columns (bottom) over 2005-2014 averaged over the Amazon region. Grey
dots represent yearly averages whereas the blue dots represent the wet season average.

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Acce





Figure 8: Time series of the sesquiterpene to isoprene ratio fluxes from model simulations. Dots representyearly averages from which the trend is calculated. Grey dots represent yearly averages.