The response of tropical rainforests to drought—lessons from recent research and future prospects

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Abstract

• Key message We review the recent findings on the influence of drought on tree mortality, growth or ecosystem functioning in tropical rainforests. Drought plays a major role in shaping tropical rainforests and the response mechanisms are highly diverse and complex. The numerous gaps identified here require the international scientific community to combine efforts in order to conduct comprehensive studies in tropical rainforests on the three continents. These results are essential to simulate the future of these ecosystems under diverse climate scenarios and to predict the future of the global earth carbon balance.
• Context Tropical rainforest ecosystems are characterized by high annual rainfall. Nevertheless, rainfall regularly fluctuates during the year and seasonal soil droughts do occur. Over the past decades, a number of extreme droughts have hit tropical rainforests, not only in Amazonia but also in Asia and Africa. The influence of drought events on tree mortality and growth or on ecosystem functioning (carbon and water fluxes) in tropical rainforest ecosystems has been studied intensively, but the response mechanisms are complex.
• Aims Herein, we review the recent findings related to the response of tropical forest ecosystems to seasonal and extreme droughts and the current knowledge about the future of these ecosystems.
• Results This review emphasizes the progress made over recent years and the importance of the studies conducted under extreme drought conditions or in through-fall exclusion experiments in understanding the response of these ecosystems. It also points to the great diversity and complexity of the response of tropical rainforest ecosystems to drought.
• Conclusion The numerous gaps identified here require the international scientific community to combine efforts in order to conduct comprehensive studies in tropical forest regions. These results are essential to simulate the future of these ecosystems under diverse climate scenarios and to predict the future of the global earth carbon balance.

Keywords Carbon · Climate · Drought · Global change · Growth · Mortality · Soil · Tropical · Water

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Contribution of the co-authors DB initiated this review and all authors significantly contributed to the writing.

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1 Introduction

Human-induced climate changes have begun to deeply impact boreal and temperate forest ecosystems (e.g. Ciais et al. 2005), but whether they are already affecting ecological functions in tropical rainforests is still an open question. Answering this question is one of the greatest challenges for scientists studying global change and for community or functional ecologists, since tropical ecosystems represent 34% of gross primary terrestrial productivity (Beer et al. 2010) and play a major role in carbon and water cycles at the global scale.

Tropical regions are characterized by high annual rainfall and high evapotranspiration. Nevertheless, a strong seasonality in precipitation and solar radiation levels (partly determined by cloud cover) occurs in most tropical regions around the world. These variations are partly driven by atmospheric movements related to the latitudinal changes in the inter-tropical convergence zone; they may lead to soil drought conditions. In environmental science, drought is defined as “a prolonged period of scanty rainfall”. This definition describes a “climatic” point of view, related to the amount of precipitation reaching the ground. However, depending on soil type, soil drainage conditions, evapotranspiration levels and depth of soil water extraction by roots, a period of scanty rainfall may—or may not—severely reduce the amount of soil water available to plants, animals or microbes. In this context, the literature on the effect of drought on tropical rainforest ecosystems does not systematically and precisely define what is meant by “drought”. Therefore, comparison of drought effects is not so obvious. Nevertheless, three types of drought are intensively studied in the current literature on tropical rainforests: seasonal droughts, extreme droughts and experimental droughts:

- Seasonal droughts correspond to seasonal variations in rainfall at a given site. This definition is based on long-term observations of precipitation regimes. One can expect that tropical rainforests are adapted to endure these seasonal droughts, as species have evolved under these conditions. Nevertheless, these seasonal variations influence ecosystem functioning and we thereafter review the existing knowledge on their response.

- Extreme droughts correspond to a period of abnormally low rainfall. A number of extreme drought events hit the tropical regions around the world over the past decades. Some may be explained by El Niño-Southern Oscillation (ENSO) events (1982/1983, 1986/1987, 1997/1998) (in Marengo et al. 2011): they have been associated with a rise in equatorial ocean surface temperatures and possibly also with modifications in the movement of the inter-tropical convergence zone (Li et al. 2006). However, in Amazonia, other major droughts are clearly unrelated to ENSO events (1980, 2005, 2010); they are linked to elevated surface temperatures in the Atlantic Ocean (Marengo et al. 2008, 2011). These extreme events may differ in intensity and duration and in the location and number of epicentres (see for instance the drought in 2005 vs. 2010 as illustrated in Fig. 1; Lewis et al. 2011). They nevertheless strongly influence the functioning of tropical rainforests and even endanger them. Thereafter, we use the existing knowledge on their impact to emphasize the limits to which these ecosystems may adapt.

- Experimental droughts correspond to through-fall exclusion experiments (Fig. 2). A given percentage of precipitation can be excluded from a given zone, and local consequences on soil, tree or ecosystem functioning are monitored. The major drawback of this approach is the surface of the considered zone: it is usually reduced (usually less than one or a few hectares). Several experiments have yet been conducted in wet tropical or subtropical regions. A list of these experiments with corresponding references is given in Table 1. As for extreme drought events, these experiments bring highly valuable information on the mechanisms involved in the response of these ecosystems to drought.

Considering these three types of drought and their specific effects on tropical forest ecosystems, we separate in this review the effects of recurrent seasonal drought on ecosystem functioning to the ones of extreme droughts and experimental ones (cf. list in Table 2). We thus focus on (i) what can be learnt from seasonal drying and (ii) what might be the consequences of more intense and/or more frequent droughts in the future. The latter question is a burning one as global earth models have made it possible to simulate an increase in the frequency and intensity of drought events in tropical regions (e.g. Cox et al. 2004, 2013; Poulter et al. 2010). The earliest simulations were highly criticized and Steinkamp and Hickler (2015), based on a global modelling analysis, concluded that there has not been any general drying trend or increase in extreme mortality events in tropical forests over the past decades. Yet, at least for the Amazonian region, global climate models converge to simulate a change in rainfall regime in this region and drier conditions over the twenty-first century (Malhi et al. 2009). Our ability to evaluate the impact of climatic change on the functioning of tropical rainforest ecosystems in the future will depend on observational, experimental and modelling approaches which include extreme drought conditions.

In this context, a considerable effort is being made worldwide to better understand the effect of drought on carbon and water cycles in tropical ecosystems at different spatial (forest stand to region) or temporal (seasonal to yearly) scales. Major research programmes have been and are being conducted in Central and South America, Asia and Africa. Within this framework, numerous new results have been published over the past 5 years. Hereafter, we provide a comprehensive
review of these recent studies and discuss whether we are currently able to evaluate the response of tropical rainforests to the future climate conditions in these regions. We also synthetize the existing knowledge on the mechanisms leading to these effects. Considerable knowledge has been accumulated on the physiological attributes of tropical rainforest tree species and their ability to support drought events. A review of this present knowledge would be highly valuable, but to keep the scope of this article within acceptable limits, we concentrate on community- and ecosystem-level responses to drought, rather than on tree- or species-level responses. We include 172 papers in this review, among which 88 were published over the past 5 years (2010–2015).

2 The influence of drought on tropical rainforest ecosystems

2.1 Tree mortality

As pointed out in Table 2, we were not able to find any information in the literature on the seasonal variations in mature tree mortality in tropical rainforest ecosystems. The main reason is that (i) tree mortality under non-exceptional years randomly occurs among seasons and (ii) seasonal drought-induced mortality is hard to detect given that mortality is a punctual and rare phenomenon difficult to catch and model apart from exceptional events (Aubry-Kientz et al. 2013). This
pattern is also consistent with the hypothesis that tropical rainforests are well-adapted to recurrent seasonal variations in soil water availability.

In contrast, the extreme drought that occurred in Amazonia in 2005 clearly resulted in abnormally high tree mortality rates (Phillips et al. 2009, 2010). Similar major die-off events were observed in Amazonia following an ENSO event (Williamson et al. 2000), in Panama (Condit et al. 1995), in tropical rainforests in China (2009–2010, 75 % increase in mortality rate; Tan et al. 2013) and in East Kalimantan on Borneo Island following an ENSO event (10–20 % increase in mortality rate; Slik 2004). Furthermore, in two sites in Amazonia (Caxiuanã and Tapajós) on long-term monitoring plots with through-fall exclusion, tree mortality also strongly increased (100 %) a few years after the beginning of the experiment compared to controls (da Costa et al. 2010). In contrast, no clear change in mortality rates was observed in Central Kalimantan, Indonesia, after the 1997 drought (Nishimua et al. 2007). Increased mortality rates in extreme drought conditions as compared to recurrent seasonal droughts clearly point out to the limits that these ecosystems are able to support. These extreme drought events or experimental studies bring extremely valuable information to where these ecosystems can be pushed out by limited water conditions.

Comparing mortality rates after extreme drought events among sites distributed worldwide is perilous because climate conditions, species composition, soil conditions, history and drought severity differ. Nevertheless, Phillips et al. (2010) concluded that for comparable drought intensities, mortality rates in Asian tropical forests were stronger than in the Amazon. More intensive research programmes should be implemented so as to confirm such spatial trends and improve ecosystem and earth model simulations, particularly in poorly studied tropical rainforests like the ones in Africa and Asia: several

| Table 1 | List of through-fall exclusion experiments conducted in wet tropical or subtropical regions |
|---------|---------------------------------|
| Site    | Location | Climate                | Period      | Reference                                  |
| Daintree rainforest | Australia | Seasonal wet tropical | 2013–       | http://www.jcu.edu.au/daintree             |
| Caxiuanã National Forest Reserve | Brazil | Seasonal wet tropical | 2002–       | Fisher et al. (2007)                       |
| Tapajós National Forest | Brazil | Seasonal wet tropical | 2000–       | Nepstad et al. (2002)                      |
| Lore Lindu National Park | Indonesia | Asesonal wet tropical | 2007–2009 | Van Straaten et al. (2011)                 |
| Fazenda Vitória | Brazil | Seasonal wet tropical | 1993–1996 | Cattánio et al. (2002)                     |
| Luquillo Experimental Forest | Puerto Rico | Asesonal wet subtropical | 2009 | Wood and Silver (2012)                     |
| Golfo Dulce Forest Reserve | Costa Rica | Seasonal wet tropical | 2007–2008 | Cleveland et al. (2010)                    |

| Table 2 | List of tree- or ecosystem-level parameters included in this synthesis and summary results from the existing knowledge under experimental droughts, seasonal droughts or extreme drought events |
|---------|--------------------------------------------------|
| Parameter | Summary results from |
| Experimental drought | Seasonal drought | Extreme drought |
| Mortality | Increase (especially large trees) | Increase likely due to change in C allocation | Increase (especially large trees) |
| Tree growth | Large trees and high-growth rate trees negatively impacted | Contrasted patterns | Reduced or even suppressed |
| Net ecosystem exchange | | | Higher C uptake |
| Gross primary productivity | | | Moderate decrease |
| Ecosystem respiration | | | Strong decrease |
| Fluxes of greenhouse gases other than CO₂ | | | Negate the positive fertilization effect of atmospheric CO₂ |
| Carbon balance | | | |
| Transpiration | Decrease through regulation of stomatal conductance | Decrease through regulation of stomatal conductance | Shift to more xeric forest types |
| Shifts in species composition | Variability in mortality rates within phylogeny | | Increase in lianas? |
| Phenology/litterfall/LAI | Slight decline in litterfall | Leaf renewal driven by solar radiation | Increase in LAI |
| Remote sensing approaches | Green-up in dry season when water is not limited | Contrasted patterns | Increase (especially in logged and disturbed forests) |
recent programmes funded by the European Commission (CARBOAFRICA, http://www.carboafrica.net) and by the National Science Foundation of China are currently making significant contributions to this much-needed research.

What mechanisms might explain the higher mortality events following extreme droughts? This question does not apply solely to tropical forest ecosystems, and a large and recent literature for various situations around the world suggests that both carbon starvation and hydraulic failure processes are involved (e.g. Hartmann et al. 2015). The “carbon starvation hypothesis” was developed by McDowell et al. (2008) (see also the recent paper of McDowell et al. (2013) and Sevanto et al. 2014) and states that tree mortality occurs mainly because reduced photosynthesis during drought periods limits the accumulation of non-structural carbohydrate compounds in trees. In contrast, hydraulic failure occurs when xylem vessels are embolized. Several studies on tropical rainforests have clearly shown strong limitation of water transported through the xylem during dry periods (e.g. Sobrado 1997; Lopez et al. 2005; Fisher et al. 2006, 2007), thus supporting the hydraulic failure hypothesis. In contrast, Metcalfe et al. (2010b) suggest that the higher mortality in through-fall exclusion stands could be explained by carbon starvation. It is now well-known that these processes represent synergistic responses to drought, but the main issue now is to understand how these processes are linked. Because the methodologies used to assess the mechanisms causing tree mortality are complex and highly time-consuming and rely on the occurrence of major droughts, we still lack enough data to decipher and understand the origin of tropical tree mortality. This limits our ability to simulate the future of these ecosystems.

It is also important to note that several studies have reported that the increase in mortality in both through-fall exclusion studies (da Costa et al. 2010) and under natural conditions (Phillips et al. 2009, 2010) was higher for large trees than for small ones. This pattern was observed in Amazonia and Borneo (Phillips et al. 2010), but not in Asian forests (Aiba and Kitayama 2002; Itoh et al. 2012a, b). Higher mortality for large trees may be related to higher evaporative demand (higher vapour pressure deficit at the top of the canopy) (Saatchi et al. 2013) or to a higher risk of hydraulic failure because of tree height. Some other studies also reported slightly higher mortality rates for trees with low density wood than for those with higher density wood (Phillips et al. 2010; Aubry-Kientz et al. 2015). Part of these differences could be linked to differences in hydraulic failure among trees, as wood density is linked to xylem structure.

It is also important to remember that extreme mortality events can also have positive feedbacks on ecosystem dynamics and functioning. Standing dead trees and/or large fallen trees modify the structure of the canopy, increase canopy openness and enhance radiation within the canopy and on the ground..., thus changing local environmental conditions (Condit et al. 1996; Holmgren et al. 2001; Slik 2004). These conditions favour the establishment of fast growing trees and the growth of neighbouring trees, thus locally contributing to increased photosynthesis and ecosystem carbon uptake. Nevertheless, fast growing trees are characterized by low wood density and are likely to be more prone to embolism (e.g. Hacke et al. 2001). In the mid to long term, these changes may thus negatively influence the carbon balance of these forests. Whether the recent demonstration by Brien et al. (2015) that the biomass dynamics of the Amazon rainforest over the past three decades show a long-term decreasing trend of carbon accumulation is consistent with this pattern is a worthy question.

2.2 Tree growth

Strong seasonal variations in tree growth (diameter increment) in tropical rainforests have been observed (Tian et al. 1998; Baker et al. 2003; Nepstad et al. 2004; da Costa et al. 2010; Stahl et al. 2010; Grogan and Schulze 2012; Wagner et al. 2012, 2013, 2014): a decrease in tree growth is usually observed during drought periods in inventory plots, followed by a strong increase at the onset of the rainy season. Extreme droughts reduce tropical forest carbon storage by decreasing or even suppressing tree growth as compared to years with lower soil moisture deficits. Phillips et al. (2009) concluded that the 2005 extreme drought event in Amazonia halted the long-term increase in aboveground biomass (since the 1950s) in a significant number of study plots. It may be expected that water-stress-induced variation in tree growth response may be dependent to topographic position, as suggested by indirect observations (Silva et al. 2013), but there is a lack of evidence on that point.

The causes for a decline in tree growth under drought conditions are multiple:

(i) Firstly, a methodological remark must be made. Growth measurements based on dendrometer bands installed around tree trunks must be used with caution, since the bands react to bark thickness and may shrink during dry periods (because of a drier atmospheric air, e.g. Stahl et al. 2010). A decrease in trunk circumference during dry periods, or at least a decrease in circumference increment, could therefore be due to the shrinkage alone, or to a “real” decrease in tree growth (cell division in cambial tissues), or both. This widely used methodology has proven efficient to study long-term tree growth, but short-term variations (at seasonal scale) can lead to misinterpretation. Chitra-Tarak et al. (2015) have recently published a novel method to take this bias into account.

(ii) Soil water has been found to be a predominant driver of seasonal variations in tree growth. Reduced relative
extractable water during drought periods was found to be the main environmental factors causing temporal variations in tree growth in an extensive study on 205 trees in French Guiana (Wagner et al. 2011). Precipitation and the duration of the drought season were the best predictors of tree growth variations in an African study (Ouédraogo et al. 2013). This was also the case in an overview across tropical regions (Wagner et al. 2014).

(iii) Third, solar radiation and daily minimum air temperature also contribute to seasonal variations in tree growth, though to a lesser extent than soil water availability (Clark et al. 2010; Wagner et al. 2012). The air temperature effect may be related to the limiting effect of high temperatures on photosynthesis. Interpreting the radiation effect is complex, as radiation may indeed influence photosynthesis levels. Yet, photosynthesis and growth are not directly linked: non-structural carbohydrates stored in trees also contribute to secondary growth, with various time lags (days to months) (Richardson et al. 2013).

The physiological mechanisms leading to seasonal variations in tree growth are not fully understood either. Secondary growth is known to decrease during drought periods when xylem embolism occurs, resulting in the disruption of water columns in the trunk and in branches and in the reduction of water supply to leaves (Tyree and Cochard 1996). The role of non-structural carbohydrate reserves should be considered too: these reserves may not only play a major role in explaining seedling survival during drought periods (Obrien et al. 2003; Kruijt et al. 2004). They also arise because of discrepancies in seasonal variations in carbon fluxes (Fig.3). Gross primary productivity (GPP) and ecosystem respiration (ER)—the two CO2 fluxes that characterize the main exchange in carbon between the ecosystem and the atmosphere—usually decrease during seasonal droughts (see van der Molen et al. 2011 and Shi et al. 2014 for a review). This decrease is largely explained by physiological [down-regulation of leaf, trunk or root gas exchange; decrease in heterotrophic respiration] or structural [leaf senescence, leaf shed] changes.

One major result obtained thanks to these studies is that differential changes in ecosystem photosynthesis (GPP) and respiration (ER) during dry periods induce large seasonal variations in the ecosystem carbon balance, as determined by net ecosystem carbon exchange (NEE). NEE is calculated as the algebraic difference between ER and GPP [the ecosystem is considered to be a carbon sink when NEE is negative]. Large discrepancies in seasonal variations in NEE across tropical rainforests have been found. Some studies showed no clear effect of seasonal soil drought on NEE (Carswell et al. 2002). In contrast, other studies showed a decrease in carbon uptake during dry periods (Williams et al. 1998; Keller et al. 2004), whereas others showed higher [more negative] NEE values during dry seasons than during wet seasons (Saleska et al. 2003; Goulden et al. 2004; Baker et al. 2008; Bonal et al. 2008; Yan et al. 2013). At the latter sites, tropical rainforest ecosystems accumulate more carbon when water is a limiting factor than during the wet seasons. Some of these discrepancies could be related to the overestimation of ER in early studies that used the eddy covariance method (see Saleska et al. 2003; Kruijt et al. 2004). They also arise because of differences in species composition, soil conditions and partly also to drought stress levels reached at the sites.

2.3 Carbon fluxes

An important consequence of higher mortality rates during broad-scale mortality events is the reduced capacity of impacted forests to absorb CO2 and the huge amounts of carbon that can be released into the atmosphere as the dead trees decomposed. This decomposition is slow (one to several decades) and varies widely according to wood density and branch or trunk size (Hérault et al. 2010). Direct, long-term measurements of carbon fluxes are recorded at various eddy flux tower sites in tropical rainforest ecosystems (now in Amazonia, Africa and Asia) and allow to characterize seasonal variations in carbon fluxes (Fig. 3). Gross primary productivity (GPP) and ecosystem respiration (ER)—the two CO2 fluxes that characterize the main exchange in carbon between the ecosystem and the atmosphere—usually decrease during seasonal droughts (see van der Molen et al. 2011 and Shi et al. 2014 for a review). This decrease is largely explained by physiological [down-regulation of leaf, trunk or root gas exchange; decrease in heterotrophic respiration] or structural [leaf senescence, leaf shed] changes.

Interestingly, drought does not influence growth in trees of different sizes or with differing growth rates in a similar way: long-term drought in through-fall exclusion experiments mainly affected the growth of large trees, whereas the small trees were barely impacted (da Costa et al. 2010). Similar trends were observed in natural forests in Amazonia and Africa (Ouédraogo et al. 2013; Phillips et al. 2009; Wagner et al. 2013, 2014). Furthermore, trees with high radial growth rates under moist conditions showed the strongest relative decrease under through-fall conditions in Indonesia (Moser et al. 2014). These effects are not intuitive since reduced soil water availability necessarily displays a vertical gradient: the top layers are usually drier than the deeper ones. One could therefore expect that trees with superficial roots would be affected the most. Yet, Stahl et al. (2013b) demonstrated that there is no clear relationship between tree size and soil water extraction depth during the dry season: small trees are also able to use water from deep soil layers. Other explanations may have to do with ontogenetic shifts in drought-adaptive capacities (Cavender-Bares and Bazzaz 2000) or in hydraulic failure/carbon allocation strategies.
Under extreme drought conditions, GPP is generally moderately further impacted by these extreme conditions, while the decrease in ER is usually much stronger than under recurrent seasonal droughts (synthesised in Shi et al. 2014). This suggests that drought severity could be driving the relative decrease in GPP and ER in these ecosystems. If this is verified, such patterns should be included in modelling approaches. A question which merits further attention then arises: what is the future of the carbon that is not released during severe dry periods, as compared to moderate dry ones? Is this carbon directly released to the atmosphere once heavy rainfalls occur or is it, partly or completely, stored in the ground?

The following mechanisms have been identified as contributing to the GPP and ER pattern under seasonal or extreme droughts:

(i) Tree root access to deep soil layers (Nepstad et al. 1994; Markewitz et al. 2010; Stahl et al. 2013b) limits the negative effect of reduced rainfall on photosynthetic activity. Indeed, though the root biomass of tropical rainforest trees is concentrated in the upper soil layers (Nepstad et al. 1994), deep roots still play a crucial role in the water and carbon cycles in these ecosystems (Nepstad et al. 1994; Stahl et al. 2013b). When soil water becomes limited in the upper soil layers, deep roots may provide water to the trees and thus help to prevent, or at least delay, the down-regulation of carbon and water fluxes. Reduced rainfall conditions over several weeks or even months do not always result in an instantaneous lack of water for deep-rooted trees; delayed drought effects may be observed. The GPP of deep-rooted trees may therefore remain high during drought events.

(ii) An increase in GPP could be expected during dry periods because of the production of new leaves (Restrepo-Coupe et al. 2013) and increased radiation [reduced cloud cover] as compared to wet periods (Huete et al. 2006; Brando et al. 2010; Samanta et al. 2012). The negative effect of reduced solar radiation on tree photosynthesis and growth during the wet season was experimentally demonstrated by Graham et al. (2003). This argument was also used by Condit et al. (2004) to explain high tree growth rates during the 1997–1998 ENSO event. However, this potential increase may be counterbalanced to some extent by the negative effect of drought on photosynthesis for the shallow-rooted trees when the dry season is moderate, or for most trees if the decrease in soil water availability is severe. The opposite trend between these different effects explains the complexity of the seasonal variations in GPP.

(iii) ER is the sum of the CO₂ efflux from all the ecosystem compartments. One can expect a physiologically driven decrease in cell respiration under drought (see Atkin and Macherel 2009 for a review), but contrasting results have been published for tropical rainforest trees. A decrease in the CO₂ efflux from the leaves, trunks and coarse woody debris was found in several studies (Chambers et al. 2000, 2001; Nepstad et al. 2002; Stahl et al. 2011, 2013a; Rowland et al. 2013, 2014). However, others showed slightly higher autotrophic (root and leaf) CO₂ fluxes under dry conditions (Metcalfe et al. 2010a, b; da Costa et al. 2014). Similarly, contrasting changes in soil CO₂ efflux have been found: a decrease in total soil respiration was observed in numerous studies (Araujo et al. 2002; Goulden et al. 2004; Hutry et al. 2007; Bonal et al. 2008). However, though some through-fall exclusion manipulation experiments confirmed this decrease (Sotta et al. 2007; Van Straaten et al. 2011; Wood and Silver 2012), other studies showed no effect of water exclusion on heterotrophic soil respiration (Davidson et al. 2008) and one even found an increase (total soil respiration; Cleveland et al. 2010). These discrepancies could be related to differences in soil type and potential local ecosystem and carbon cycling adaptations to climatic conditions (Cleveland et al. 2010). However, the levels of drought stress reached at the study sites (seasonal vs. extreme) and the duration of the drought do not seem to

Fig. 3 The 55-m high eddy flux tower set up in 2003 in French Guiana (Bonal et al. 2008) (credits D. Bonal). Equipment installed on top of the tower (above the canopy) allows to continuously monitor the gas exchange (CO₂, H₂O) between the atmosphere and the considered ecosystem (around 50–100 ha)
contribute to explain these differences. These results underline the complexity of the carbon cycle in tropical ecosystems submitted to drought and make it difficult to simulate these response patterns with ecosystem or earth models.

2.4 Transpiration

Tropical rainforests play a major role in the global climate system because they exchange huge amounts of water and energy with the atmosphere. Droughts of varying intensity modify the hydrological cycle over these ecosystems (Gloor et al. 2013). When tropical rainforests suffer from seasonal drought, a decrease in tree- and ecosystem-level evapotranspiration is observed (Bonal et al. 2000; Stahl et al. 2013a). When the drought conditions become harsher, the decrease in tree transpiration is much stronger, and Fisher et al. (2007) concluded that tree water use was restricted to 20 % of that in the control plots in a through-fall exclusion experiment. However, it is important to keep in mind that a decrease in evapotranspiration during a drought period actually reflects the combination of two opposite effects. First, high solar radiation during dry periods induces high evaporative demand [drier and warmer atmospheric air] that should theoretically lead to higher evapotranspiration rates. As an example, Fisher et al. (2007) confirmed higher transpiration during the dry season than during the wet season in Brazil. Second, a decrease in soil water availability leads to the down-regulation of stomatal conductance and thus to lower tree transpiration (Bonal et al. 2000; Stahl et al. 2013a). This generally observed decrease in transpiration during droughts suggests a predominant down-regulation of transpiration (reduction in soil-to-root conductance and water transport) when soil water availability is strongly limited. Nevertheless, the effect of soil water drying and strong evaporative demand on stomatal regulation can be diverging (Tardieu and Simonneau 1998; Tardieu et al. 2010). The complexity of these effects should not be overlooked when investigating observed patterns, particularly in modelling analyses.

3 What faces tropical rainforests in the future?

3.1 Species composition

Based on biogeographical, taxonomic or molecular biology studies, it has now been widely accepted that the tree species composition in tropical rainforests evolved during the Late Pleistocene and Holocene periods in conjunction with the occurrence of drier and cooler periods (Mayle et al. 2000). Even though major parts of tropical rainforests may have been remarkably resilient to dry climatic conditions (Mayle and Power 2008), the current species composition of these forests is directly linked to past climate changes (Liu and Colinvaux 1985).

Just as the tropical rainforest communities of today were shaped by past climate changes, so will tropical species distribution, community composition and diversity be altered if climate conditions drastically change in the near future (Slik 2004; Engelbrecht et al. 2007; da Costa et al. 2010). Malhi et al. (2009) simulated that future climate conditions over Amazonia will be favourable for seasonal forests rather than for dry tropical forest or dry savannahs. In contrast, some global vegetation models predict a dramatic shift of the biome in Amazonia towards dry forests or even woodland savannahs (Cox et al. 2004; Chave 2014). These predictions are somehow catastrophic considering the known resilience of these ecosystems to drought events during the Holocene (Mayle and Power 2008). Nevertheless, they are supported by a few observations in tropical rainforests where local species composition and community functioning appear to shift over two to three decades towards more xeric types, if soil water conditions are frequently limited (e.g. Condit 1998; Enquist and Enquist 2010). This shift is associated with the contrasting vulnerability of tropical species to drought: for certain taxa, or size classes, or growth characteristics (fast vs. slow growing), or successional status (early vs. late stage), some species are subject to disproportionate mortality. Through-fall exclusion experiments have also revealed a strong variability among tree species, or tree genii, in their mortality rates (da Costa et al. 2010; Nepstad et al. 2007). A precise description of the functional characteristics of tropical tree species and their physiological and morphological response to drought is far from being complete. Nevertheless, important screenings of traits related to drought have been conducted over the past decades (e.g. Bonal et al. 2000; Engelbrecht et al. 2002, 2007; Engelbrecht and Kursar 2003; Bartlett et al. 2012; Fortunel et al. 2014). The vulnerability of tropical tree species to prolonged droughts diverges and is related to intrinsic differences in various leaf (e.g. turgor loss point, osmotic potential, stomatal closure, leaf thickness, specific leaf area, water use efficiency) and wood biological properties (e.g. xylem vessel size, wood specific gravity). Further comprehensive studies on the functional characteristics of tree species in Amazonia, Africa and Asia are still required.

As an example of shift in species composition, an increase in lianas, both in terms of individuals and biomass, has been documented over the past decade in tropical rainforests (see Schnitzer and Bongers 2011 for an overview). This change in distribution may have various origins (increasing forest disturbance, changes in land use, forest fragmentation...), but it has also been suggested that climate changes, along with extreme drought events, could be playing a major role. This remains to be proven; observations are recent and correlative changes are under study. Nevertheless, lianas are known to
benefit, to the detriment of tree species, from drier air (increased evaporative demand) or soils (decreased soil water availability) (Schnitzer 2005; Schnitzer and Bongers 2011). Climatic conditions in the future may continue to be favourable to lianas and allow them to colonize new tropical rainforests, or at least to accentuate their contribution to ecosystem biomass and productivity.

These functional considerations should not hide one very important aspect in community ecology: tropical rainforests are the most diverse forests in the world (at least in terms of tree species). For a large range of ecosystem services, more diverse forest ecosystems perform better that pure ones because of positive interactions among tree species for resource acquisition (facilitation or niche differentiation mechanisms; see Cardinale et al. 2012 for a review). Basic theoretical ecology also suggests that biodiversity insures ecosystems against alteration of ecosystem functions, the insurance hypothesis (Yachi and Loreau 1999). Numerous species with a broad range of functional characteristics provide greater ecological overlaps so that if some species are loss, the remaining ones maintain similar levels of ecosystem functioning. In contrast, Mouillot et al. (2013) recently demonstrated the major role of rare species and functional vulnerability in the stability of ecosystem functioning. Unfortunately, very few studies on the relationship between biodiversity and ecosystem functioning have been conducted in tropical regions and those few mainly concern experimental plantations (e.g. Potvin and Gotelli 2008; Forrester 2014). Tropical rainforests with different levels of species diversity or with different species composition may respond differently to future droughts. A completely new area in ecological research is needed to answer this question.

### 3.2 Phenology patterns

Tropical rainforests are composed of species that cover the full range of leaf phenology patterns (i.e. emergence and fall), from strict deciduousness to evergreen (Loubry 1994; Viennois et al. 2013). The leaf area index (LAI) in these ecosystems remains high all year long and they are therefore classified as evergreen forests. Nevertheless, inconsistent seasonal patterns in LAI have been observed in Amazonia. Malhado et al. (2009) and Wagner et al. (2013) observe no seasonal change in LAI, whereas several studies conclude to clear seasonal dynamics in LAI or litterfall (e.g. Juarez et al. 2009). A peak in litterfall is usually observed during the early stages of the dry season in Amazonia (see Chave et al. 2010 for an overview). The origin of this later pattern remains unexplained: reduced water availability does not seem to provide an explanation (Wright and Cornejo 1990; Xiao et al. 2006; Miranda et al. 2011). Seasonal variations in leaf phenology were found to be driven by solar irradiance (Wright and van Schaik 1994; Xiao et al. 2006; Wagner et al. 2013). The peak in leaf fall coincides with a peak in radiation (Wagner et al. 2013) and is associated with a greater emergence of new leaves at the onset and during the dry season, a period when cloud cover is usually low or absent and solar radiation is therefore strong. Kim et al. (2012) showed that including a light-controlled phenology module into a terrestrial biosphere model (ED2 model) strongly improved its ability to capture seasonal patterns in net ecosystem productivity and litterfall fluxes. The greater proportion of young leaves during the dry season could enhance carbon assimilation during the dry season and at the onset of the wet season, at least as long as stomatal regulation related to drought stress does not limit photosynthesis. Indeed, young leaves are characterized by higher photosynthetic capacity than mature or senescent ones. However, this argument has been contradicted by studies showing a delayed development of the photosynthetic apparatus in young tropical rainforest leaves and lower photosynthetic efficiency compared to mature ones (Kursar and Coley 1992).

When considering the influence of extreme drought events, strong changes in ecosystem leaf area were indeed observed (Hilker et al. 2014). However, interestingly, long-term through-fall experiments showed no clear impact of severe drought on litterfall over the first and second year (Brando et al. 2008); a slight decline in litterfall compared to control plots occurred only at the third year. This pattern confirms the drought resistance of tropical ecosystems when an extreme event does not last too long, but also the limits to this resistance.

Optical remote sensing has been widely used over the past decade to analyse the effect of extreme drought on ecosystem phenology at various time scales and over large areas. After the discovery—using satellite measurements—of an unexpected canopy green-up during the dry season in Amazonia during the 2000–2005 period (Huete et al. 2006), Saleska et al. (2007) detected an increase of this green-up during the extreme dry season in 2005. This green-up was interpreted as being consistent with the leaf flush usually observed during dry conditions. However, Samanta et al. (2010)—and more recently Morton et al. (2014)—argued that these conclusions were based on irreproducible data and potential artefacts: the satellite data quality [partly due to cloud cover] may not have been good enough to support such a strong interpretation. The authors then concluded that no increase in green-up actually occurred during extreme drought events, a result consistent with the decline in greenness (less photosynthesis) during the 2010 drought event over Amazonia (Xu et al. 2011). However, an increase in greenness was recently observed by Maeda et al. (2014) during the dry season, even after correcting for all potential sun/sensor geometry artefacts. These results are clearly contradictory and a recent study tempted to reconcile these diverging views (Hilker et al. 2014). These authors conclude that the new, better quality satellite data such as those obtained with C5-MODIS...
eliminate such potential artefacts and that the occurrence of canopy cover green-up during seasonal dry periods, or extreme ones, depends on whether trees can access to deep soil water. As a result, some parts of the Amazon forest do show green-up during the dry season, though limited, since, during the cloudy wet season, the reduced light limits ecosystem photosynthesis. However, in other parts of the Amazon forest where tree access to soil water is severely limited, no green-up occurs; a decline in greenness may even be observed via satellite. Studying tropical forest phenology with satellites data is thus feasible, but precisely assessing the impact of severe drought events is not yet fully solved (Hilker et al. 2014).

3.3 Carbon balance

Considerable literature over the past decade has demonstrated the positive role of tropical rainforests in capturing large amounts of atmospheric CO$_2$ (e.g. Baker et al. 2004; Malhi et al. 2004; Ichii et al. 2005; Lewis et al. 2009; Phillips et al. 2009; Pan et al. 2011). Net aboveground biomass gain has been confirmed on numerous inventory plots in Amazonia or Africa: Phillips and Lewis (2014) estimate that the relative rate of increase is around 0.30 and 0.29 % for Amazonia and Africa, respectively. Accumulation of carbon in the tropical terrestrial biosphere strongly contributes to slowing the rate of increase of CO$_2$ into the atmosphere, thus reducing the greenhouse gas effect. Undisturbed tropical rainforests have been acting as a strong carbon sink in this way for decades.

The question now is whether they will be able to continue playing this role under future climate, particularly if the occurrence of extreme drought events increases. The large mortality event associated with the extreme 2005 drought in Amazonia was equivalent to several decades-worth of carbon accumulated by the tropical rainforest (Phillips et al. 2009; Gloo et al. 2012). A recent analysis of atmospheric CO$_2$ measurements concluded that the unburned part of the Amazon basin [carbon release through fires is deducted in carbon balance calculations] shifted from a net carbon sink during a wet year to a carbon-neutral status during an extreme drought year (i.e. 2010) (Gatti et al. 2014). This shift was explained not only by changes in air temperature and vapour pressure deficit, but also by reduced soil water availability. An increase in the frequency of such major drought events in the future is therefore likely to modify the positive carbon uptake role played by undisturbed tropical rainforests and may negate the positive fertilization effect of increased CO$_2$ concentrations (Cox et al. 2013). Yet, serious inconsistencies among simulation studies can be found: Phillips and Lewis (2014) reviewed the predictions concerning the carbon balance in Amazonian and African tropical rainforest ecosystems: in the future, these ecosystems may act as potential sinks, or become carbon sources, or have a neutral balance. Furthermore, a recent pan-tropical study based on 150 years of tree ring analysis showed that increased atmospheric CO$_2$ concentration induced an increase in intrinsic water use efficiency (the ratio of leaf CO$_2$ assimilation over stomatal conductance) but not an increase in individual tree growth (van der Sleen et al. 2015).

The uncertainties around these simulations are largely related both to the structure and the parameterization of the global circulation vegetation models in use today. As for phenology patterns, a major challenge for remote sensing or modelling approaches is to reconcile satellite observations with observed measurements of ecosystem carbon fluxes and storage. A wide range of ecosystem or earth models have been elaborated and tested to date (e.g. Saleska et al. 2003; Ichii et al. 2007; Baker et al. 2008; Sitch et al. 2008; Fisher et al. 2010; Sakaguchi et al. 2011; Rowland et al. 2014) and a large body of literature addresses these issues. Though we will not discuss this literature in detail here, it is evident that no model is yet able to encompass all the complex plant- and ecosystem-water interactions under limiting soil water conditions. Solid improvements have been made over the past decade in matching the more mechanistic models with local observed data. Powell et al. (2013) perfectly summarizes the current state-of-the-art: “… terrestrial biosphere models are competent at predicting plant and ecosystem carbon fluxes under the present climate, but still require substantial development for predicting the consequences of severe drought scenarios.”.

The influence of drought on the exchange of greenhouse gases other than CO$_2$ between tropical rainforests and the atmosphere is rather unknown. A few studies conducted over the past decade showed significant seasonal variations in N$_2$O or CH$_4$ fluxes (Davidson et al. 2008; Itoh et al. 2012a, b; Wood and Silver 2012; Yan et al. 2014), but further comprehensive studies are required in order to include the effect of drought on these fluxes in ecosystem or earth models.

3.4 Fires

A major risk of future droughts—and one that may radically change species composition—is forest fire. Natural forest fires occasionally occurred in tropical rainforests throughout the Holocene period (Saldrarriaga and West 1986) and contributed to shape the biogeography of the actual tropical rainforests. Nevertheless, the major drought events of the past decades have resulted in an increased number of major forest fires in tropical rainforests (Cochrane et al. 1999; Alencar et al. 2006; Aragão et al. 2008). The risk of major fires is also expected to increase in the twenty-first century (Davidson et al. 2012). These fires have altered species composition and forest dynamics in large areas and will have long-term consequences on these ecosystems (Brando et al. 2014). Some evidence
suggests that in tropical rainforests, selective logging (Siegert et al. 2001), pasture burning (Nepstad et al. 1999) and forest fragmentation (Cochrane and Barber 2009) lead to an increased susceptibility of the forest canopy to fire. This increase is due to a greater amount of dead wood in these forests, a lower relative humidity in the understory which increases the drying speed of the dead wood and a greater penetration of forests by human activities. For example, opening of the canopy by logging generates faster air movements in the canopy so that the increase in fire-line intensity in logged forests in the Amazon region is estimated to be twice as intense as in intact forests (Cochrane 2003). Furthermore, drought-related forest fires obviously release huge amounts of carbon into the atmosphere (Grace et al. 2014); at the regional scale, Gatti et al. (2014) showed that forest fires occurring during the extreme 2010 drought contributed to shift the annual carbon balance of the Amazon basin from a carbon-neutral status to a strong carbon release zone (0.48±0.18 Pg of carbon per year). In addition to the direct effect of fires on carbon balance, carbon losses from tree mortality in fire-affected forests have been shown to be cumulative through time, with an increase in large tree mortality three years after the fire event (Barlow et al. 2003).

4 Conclusions and research gaps

As synthesized here, a huge effort has been made over the past decade to improve our understanding of the response of tropical rainforest ecosystems to seasonal variations in precipitation and to extreme soil water conditions. The existing literature points to the adaptive capacity of these ecosystems to support seasonal droughts, but also to the great diversity and complexity of this response. The recent extreme drought events that occurred in these regions, the studies conducted in experimental droughts and the ongoing modelling studies allowed to greatly improve our knowledge when these ecosystems are pushed out of the soil water conditions during recurrent seasonal droughts. We now have better ideas about the potential limits for the carbon and water cycle and balance of these ecosystems to support extreme drought conditions. Our ability to model this response and simulate the potential consequences of future climate conditions has thus greatly improved, but large uncertainties, and even some discrepancies, still exist. This effort should thus be continuously supported, considering the major role played by these ecosystems at the global scale.

We have identified a number of research gaps to fill in the current knowledge on the response of these ecosystems to severe drought conditions. We highlight the most important below. These gaps require the international scientific community to combine efforts in order to conduct comprehensive studies in tropical rainforests on the three continents. These results are essential not only to simulate the future of these ecosystems but also to predict the future of the global earth carbon balance.

4.1 Taking combined stresses into account

An important point to consider is that the increase in the intensity and frequency of drought events will be accompanied by other changes in environmental conditions, such as increases in temperatures, vapour pressure deficit or atmospheric CO₂ concentrations. This review specifically targeted the effects of soil drought, but one must not lose sight of the consequences of other changes and of the potential feedback controls (Malhi et al. 2009; Cernusak et al. 2013). Niinemets (2010) concluded that tree response to combined stresses, though none of them may be extreme separately, can be much greater than tree response to one single extreme stress. A concerted research effort should therefore be made in the near future to study combined effects. Modelling studies or large-scale manipulation studies combining soil drought, increase in CO₂ concentrations and temperature, or combining soil drought and nutrient levels should be encouraged in these forests to test the effect of these combined changes (Cavaleri et al. 2015).

4.2 Relating species-level and ecosystem-level responses to extreme droughts

Though we must maintain our research efforts, our understanding of tropical rainforest species vulnerability to drought has improved over the past decade. Ecosystem functioning depends on species composition and ecosystem response to drought results from each individual species response to drought. However, these responses do also depend on interactions among species (competition, facilitation), on interactions between species and their attendant microbial communities and on changes in species composition. Information about these interactions and changes is totally lacking. Manipulating species composition on large scales in these forests seems unrealistic, but building new models combining mechanistic and community dynamics approaches should help us understand the effects of climate changes on ecosystem functioning.

4.3 Understanding the role of microbial communities

There is evidence that plant resistance to drought is improved when fungal communities densely colonize the roots (e.g. Augé 2000; overview in Lehto and Zwiezk 2011). Yet, it has been shown that fungal community structure is strongly linked to tree species composition and tree species richness in tropical rainforests (Peay et al. 2013). A change in tree species composition would thus influence fungal communities and their role in ecosystem functioning. Studies on the role of fungal communities in ecosystem functioning in tropical rainforests are very rare. A comprehensive study of these
interactions is needed in order to understand the impact of seasonal drought and extreme drought events on belowground microbial communities and clarify their role in ecosystem functioning under climate changes.

4.4 Taking repetitive drought events into account

Most observations on the effect of drought on ecosystem functioning, with but a few exceptions, are based on a single drought event. The impact of repetitive events is still far from being understood, as is the resilience of tropical ecosystems to these events. Phillips et al. (2010) pointed out that the extreme 2005 drought in Amazonia was still affecting mortality rates in these ecosystems 2 years after the drought. How well these ecosystems can adapt to more frequent severe drought events [ENSO-related or others] in the future remains unknown. Is the resistance of tropical rainforest trees weakened by a first extreme drought event which subsequently increases their vulnerability? The recent occurrence of extreme drought events at low intervals in Amazonia or Asia brings interesting opportunities to study the effect of recurrent drought events in situ.

4.5 Assessing the adaptive potential of tropical rainforest tree species to droughts

Change in species composition is one way these ecosystems can naturally adapt to severe drought conditions, but functional adaptive mechanisms can also take place. One can wonder how far such adaptive mechanisms can go and when their limits will be reached. Tree tolerance to environmental stresses increases throughout ontogeny: over the next century, trees that are young today may well prove able to adapt to upcoming climatic conditions. However, how well tropical tree species will adapt to drought over the long-term is largely unknown. Among the strategies species use to survive drought conditions, carbon allocation deserves serious studies. A major gap exists in our knowledge concerning how carbon allocation patterns change in tropical rainforest trees under varying environmental conditions, though a few recent studies have provided interesting results: a clear asynchronism between leaf production and tree growth was observed (Wagner et al. 2013). This pattern has recently been linked with a different timing in carbon allocation between compartments (Malhi et al. 2014; Rowland et al. 2014). The authors also demonstrated different allocation patterns between the above- and belowground compartments. As an example, Rowland et al. (2014) showed a shift in the allocation of photosynthetic products from foliage and wood in the wet season to fine roots in the dry season. These allocation patterns are likely complex since they undoubtedly vary with drought intensity (Branco et al. 2008; Doughty et al. 2014). Comprehensive screenings on species variability in carbon and nitrogen allocation patterns should be conducted: they should include the consequences of both recurrent seasonal variations as well as extreme drought events.

4.6 Improving the ability of ecosystem or earth models to simulate drought responses

Even though a strong effort has been made in the last decade to include as many fine-scale functional processes as possible in mechanistic forest ecosystem models or earth models (e.g. Fisher et al. 2007; Sakaguchi et al. 2011; De Weir et al. 2012; Powell et al. 2013; Rowland et al. 2014), their ability to simulate the future response of tropical ecosystems to major drought events remains inadequate. In the context of climate change, large uncertainties on future carbon stocks or fluxes persist (Poulter et al. 2010). These authors have identified research guidelines to improve the models, but to parameterize them, more observational and experimental data on the response of tropical ecosystems to drought events must be made available in concert. To improve simulations of climate change in the twenty-first century and its long-term effect on the ecosystem carbon balance, we should concentrate on phenology and mortality patterns and on CO2 and other greenhouse gas fluxes among the different compartments in the ecosystem, particularly autotrophic respiration.

4.7 Evaluating the long-term impact of punctual extreme events

Most studies published so far have addressed the short-term consequences of extreme droughts; only very few (mainly experimental through-fall studies) have attempted to evaluate their long-term impact (Fig. 4). Despite their tremendous value in the study of response mechanisms, experimental studies cannot mimic the impact of successive, high-frequency major

Fig. 4 Installation of an automatic weather station in the tropical rainforest in French Guiana (credits B. Hérault). Long-term monitoring of environmental conditions in remote forest inventory plots is a prerequisite to understand the effect of seasonal or extreme drought on tree and ecosystem functioning and simulate future changes
droughts. The recent occurrence of several droughts in Amazonia and Asia with a return time lesser than 10 years may not give the forests time to completely recover between events and thus offers a major opportunity to evaluate the long-term impact of punctual extreme events. Remote sensing approaches provide the excellent tools to analyse these effects.

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