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Seasonal controls on the exchange of carbon and water in an Amazonian rain forest

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The long-term resilience of Amazonian forests to climate changes and the fate of their large stores of organic carbon depend on the ecosystem response to climate and weather. This study presents 4 years of eddy covariance data for CO$_2$ and water fluxes in an evergreen, old-growth tropical rain forest examining the forest’s response to seasonal variations and to short-term weather anomalies. Photosynthetic efficiency declined late in the wet season, before appreciable leaf litter fall, and increased after new leaf production midway through the dry season. Rates of evapotranspiration were inelastic and did not depend on dry season precipitation. However, ecosystem respiration was inhibited by moisture limitations on heterotrophic respiration during the dry season. The annual carbon balance for this ecosystem was very close to neutral, with mean net loss of 890 ± 220 kg C ha$^{-1}$ yr$^{-1}$, and a range of −221 ± 453 (C uptake) to +2677 ± 488 (C loss) kg C ha$^{-1}$ yr$^{-1}$ over 4 years. The trend from large net carbon release in 2002 towards net carbon uptake in 2005 implies recovery from prior disturbance. The annual carbon balance was sensitive to weather anomalies, particularly the timing of the dry-to-wet season transition, reflecting modulation of light inputs and respiration processes. Canopy carbon uptake rates were largely controlled by phenology and light with virtually no indication of seasonal water limitation during the 5-month dry season, indicating ample supplies of plant-available-water and ecosystem adaptation for maximum light utilization.

1. Introduction

Tropical forests are closely coupled to climate, exerting a strong influence on temperature and precipitation patterns whilst these same weather and climate patterns dictate where particular forest types can establish and persist [Holdridge, 1947]. The interactions between regional and global climate and the Amazonian rain forest are uncertain. Both model results and field studies show wide variability in the spatial patterns and seasonality of forest growth, respiration, and water exchange [e.g., Saleska et al., 2003; Schaphoff et al., 2006]. Mechanistic understanding of the forest responses to climatic factors (particularly temperature, light, and moisture) is required to improve ecosystem process models for tropical forests and to enable more accurate projections of possible responses to changes in climate.

The Amazon Basin accounts for 50% of the world’s undisturbed tropical rain forest [FAO, 1992], 10% of global terrestrial net primary productivity [Melillo et al., 1993], and a major portion of the global surface evaporation [Choudhury and DiGirolamo, 1998]. Much of the Amazon maintains a green canopy throughout the dry season by acquiring water through deep roots [Nepstad et al., 1994] and possibly by hydraulic redistribution of water by plants [Oliveira et al., 2005]. Huete et al. [2006] found widespread ‘green-up’ of Amazonian rain forest during the dry season, with new leaf production during the period of maximum temperature, the most sunlight, and minimum precipitation.

Previous Amazonian studies have reported diverse seasonal patterns in the net ecosystem exchange of CO$_2$ from forests. Some sites found enhanced uptake of CO$_2$ during the dry season [Saleska et al., 2003; Goulden et al., 2004], others reported decreased uptake during the dry season [Malhi et al., 1998].
Araújo et al. [2002], von Randow et al. [2004], and others showed no seasonality in the exchange patterns [Carswell et al., 2002]. A data-model comparison for the Tapajós National Forest found that the Terrestrial Ecosystem Model (TEM) [Tian et al., 1998] and Integrated Biosphere Simulator (IBIS) [Botta et al., 2002] model predicted seasonality opposite to observed patterns [Saleska et al., 2003]. Net carbon uptake was observed in the dry season due to lower seasonal respiration rates [Saleska et al., 2003], whereas models predicted carbon release in the dry season due to water limitations on photosynthetic uptake of CO₂.

Similarly, evapotranspiration (ET), the combination of surface evaporation and plant transpiration, has been found to peak at some forest sites during the dry season when radiation inputs were highest [Hutyra et al., 2005; da Rocha et al., 2004; Carswell et al., 2002; von Randow et al., 2004; Shuttleworth, 1988], but at other sites maximum ET occurred during the wet season when water availability was highest [Malhi et al., 2002; Vourlitis et al., 2002]. The observed divergence between sites is likely due to differences in the actual water available to the vegetation, plus differences in phenology and radiative drivers. The amount of moisture available to a forest affects the forest's physical structure, ecophysiology, and flammability. Moisture availability is a function of not only incoming precipitation, but also the depth and texture of the soil, the depth of the water table, transpiration demands of the forest, soil capillarity, site hydrology, and the vertical distribution of roots. Drier forests can behave like moister forests if deep roots and/or favorable soils provide access to water throughout the dry season months.

Global Climate Models (GCM) generally predict decreases in Amazonian ET during the dry season, in phase with precipitation [Dickinson and Henderson-Sellers, 1988; Werth and Avissar, 2004]. Lee et al. [2005] updated the National Center for Atmospheric Research Community Atmospheric Model to include both hydraulic redistribution and deep roots in the Amazon. This model produced higher dry season ET relative to control runs, but ET still maximized during the wet season. Evidently we need a better understanding of the controls on H₂O exchange in order to improve models to predict forest flammability and to forecast the effects of drought on forest species abundances, biomass distributions, and rates of photosynthesis and ecosystem respiration.

To gain insight into the mechanisms controlling the exchange of carbon and water at the Tapajós old-growth forest, we first summarize observed local meteorology and energy exchange, and then present detailed methods, data processing techniques and validation strategies necessary for making accurate, unbiased eddy covariance measurements in a remote rain forest. We address two major scientific questions: (1) What are the controls on seasonal and inter-annual variations of net ecosystem exchange of CO₂, respiration, and photosynthesis, and water exchange? (2) Is forest growth water-limited during the dry season, or on an annual basis?

2. Methods

2.1. Site Description

Our study was part of the Brazilian-led Large-Scale Biosphere-Atmosphere Experiment in Amazonia (LBA-ECO). The site is located in the Tapajós National Forest (TNF; 54°58'S, 2°51'W, Pará, Brazil) near Km 67 of the Santarém-Cuiabá highway (BR-163). The TNF is bounded by the Tapajós River to the west and the BR-163 highway on the east, extending from 50 km to 150 km south of the city of Santarém, Pará, Brazil. East of BR-163 the landscape is extensively developed for agriculture. The tower was located ~6 km west of the BR-163 highway and ~6 km east of the Tapajós River, in an area of largely contiguous forest extending for tens of kilometers to the north and south.

The soils at this site are predominately nutrient-poor oxisols with pockets of sandy ultisols, both having low organic content and cation exchange capacity [Silver et al., 2000]. During well drilling at a nearby site with similar soils, the water table was found to be at ~100 m depth [Nepstad et al., 2002]. The forest is on flat terrain and has a closed canopy with a mean height of approximately 40–45 m and emergent trees reaching up to 55 m. There are few indications of recent anthropogenic disturbance other than small hunting trails. This forest can be classified as 'primary' with abundant large logs, numerous epiphytes, an uneven age distribution, and emergent trees [Clark, 1996]. Ground-based biometric plots were established at this site in July, 1999. See Rice et al. [2004] and Vieira et al. [2004] for more complete descriptions of the forest structure and growth dynamics.

A 64 m tower (Rohn 55G, Peoria, IL) was instrumented for eddy covariance measurements which commenced in April, 2001 and continued until the tower was destroyed when a falling tree hit the guy wires in January 2006. Three modular enclosures (approximately 1 m x 0.6 m x 0.2 m) containing all the key instruments and data loggers were mounted on the tower to keep inlet tubes short (~2 m) (Figure 1). Eddy-flux measurements were made at a height of 57.8 m with a sample rate of 8 Hz. A 3-axis sonic anemometer (CSAT-3, Campbell Scientific, Logan, UT) was mounted with the air sample inlet located 20 cm from the anemometer. The flux system drew sample air from the inlet through a 50 mm diameter Teflon filter and 9.5 mm (inner diameter) Teflon PFA tubing to a closed-path infrared gas analyzer (IRGA, LI-6262, Licor, Lincoln, NE). The eddy system sample cell (11.9 cm²) was pressure-controlled at 66.6 kPa with a mass flow rate of 6000 sccm, providing a cell-flushing time of 0.078 s. This system design maintains the advantages of the closed-path sensor (e.g., precise instrument calibration, constant pressure and temperature), while also adding some of the advantages (e.g., minimal attenuation of high-frequency fluctuations) attributed to open-path designs. This system is particularly suitable for deployment with very tall vegetation where problems accrue due to long sample-tubes from the top of the tower.

Calibrations of the eddy system for CO₂ were made every 6 hours (April 2001–November 11, 2002 and March 29, 2003–November 15, 2003) or 12 hours (November 12, 2002–March 29, 2003 and November 15, 2003–January 24, 2006) using 325, 400, and 475 ppm CO₂ standard gases traceable to world standards. The instrument was zeroed every 2 hours using a zero air generator (Parker Balston 74–5041, Haverhill, MA). The long-term accuracy of the instruments was ensured by measuring a surveillance standard (traceable to NOAA/CMDL standards at 380.45 ppm) once per week, this tank lasted through the duration of the
measurements. Calibrations for water vapor were made using the daily fluctuations of $T_v - T_k$, where $T_v$ is the sonic temperature (derived from the speed of sound provided by the sonic anemometer, closely approximating the virtual temperature) and $T_k$ is the ambient temperature. This approach was necessary due to failures in the chilled mirror hygrometers originally installed for this purpose (see the auxiliary material for additional details about calibration methods).

Vertical profiles of CO$_2$ and H$_2$O concentrations were measured at 8 levels on the tower (62.2, 50, 39.4, 28.7, 19.6, 10.4, and 0.91 m). Sample air was drawn at 1000 sccm through the 8 profile inlets in sequence (2 minutes at each level). The profile concentration data were used to estimate the change in vertical average concentration between the ground and flux measurement height in order to calculate the column average storage of CO$_2$. The profile IRGA was zeroed between each profile sequence and an absolute calibration at 325, 400, and 475 ppm was made every 6 or 12 hours, as it was for the eddy CO$_2$ measurements.

A suite of environmental measurements was also made on the tower (Table 1). Data loggers (CR-10X, Campbell Scientific, Logan, UT), controlled the overall operation of the system. The data were downloaded via coaxial cable to a computer, housed in a climate controlled hut near the tower.

### 2.3. Data Processing and Analysis

The net ecosystem exchange (NEE) of CO$_2$ between the forest and the atmosphere was computed as

$$\text{NEE} = \frac{w c'}{2} + \frac{\partial}{\partial t} \int_0^h c(z) dz$$  \hspace{1cm} (1)$$

where the first term on the right hand side is the covariance between vertical wind velocity fluctuations ($w'$) and fluctuations in the concentration of the scalar ($c'$, CO$_2$). The second term is the rate of change in the canopy storage, where $z$ is the height above the ground surface, $h$ is the flux measurement height, $t$ is time, and the overbar denotes a time average [Baldocchi et al., 1988]. The vertical coordinate for wind velocities is positive upward, thus positive values for fluxes denote emission and negative values denote uptake. Concentrations of CO$_2$ and H$_2$O were calculated using output from the IRGA's raw signal using a third order polynomial fit to the calibration data. CO$_2$ concentrations were corrected for water vapor. The temperature and pressure inside the sample cell were constant and thus no density fluctuation corrections were required (the data were represented as mole fraction in dry air [see Webb et al., 1980]).

Cospectral analyses of CO$_2$, H$_2$O, and heat flux measurements were done to assess the reliability of the flux data and to verify if appropriate averaging intervals have been used to capture all of the flux-carrying eddies [Kaimal et al., 1972]. An ogive analysis [Lee et al., 2004] provided an independent check on the adequacy of sampling intervals by looking for an asymptotic plateau in the cumulative sums of the cospectra (between 1 Hz and 34.2 minutes). The daytime ogives for CO$_2$, H$_2$O, and heat fluxes (Figure S1.) indicate that for this site a 30 minute averaging period was appropriate. We did not examine averaging intervals beyond 34.2 minutes due to the instrument calibration schedule, but the ogives indicate that the low frequency fluxes were adequately captured. There was some attenuation of high frequency (above 0.1 Hz) components of the water vapor flux due to adsorption and desorption along the sample tube walls and inlet filters, but attenuation losses were low (<2%) because of the short sample tube lengths. We corrected for...
Table 1. List of Environmental Measurements, Instruments, and Measurement Heights on the Tower

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Instrument</th>
<th>Height on Tower</th>
</tr>
</thead>
<tbody>
<tr>
<td>Net radiation</td>
<td>Rebs Q7.1 with RV2 ventilation</td>
<td>64.1 m</td>
</tr>
<tr>
<td>Photosynthetically active radiation (PAR)</td>
<td>Licor 190-2A</td>
<td>63.6 m and 151.4 m</td>
</tr>
<tr>
<td>Aspirated air temperature</td>
<td>Spinning cup anemometer, Met One 010C</td>
<td>57.9 m</td>
</tr>
<tr>
<td>Atmospheric pressure</td>
<td>MKS 627A Baratron pressure transducer</td>
<td>Ground-level</td>
</tr>
<tr>
<td>Dew point hygrometers</td>
<td>EdgeTech 200M</td>
<td>57.9 m</td>
</tr>
<tr>
<td>Wind speed</td>
<td>Spinning cup anemometer, Met One 010C</td>
<td>64.1 m</td>
</tr>
<tr>
<td>Wind direction</td>
<td>Met One 020C</td>
<td>42.6 m</td>
</tr>
<tr>
<td>Precipitation</td>
<td>Texas Electronics 076B-4</td>
<td>42.6 m</td>
</tr>
</tbody>
</table>

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reported errors are 95% confidence intervals calculated by bootstrapping the error distributions during similar (e.g., season, hour, PAR level) time periods [Richardson and Hollinger, 2005]. Seasonal mean results are based on the mean dry season interval extending from July 15–December 15 with the remainder of the year being considered the wet season.

3. Results

3.1. Weather and Climate

[23] Tables 2 and 3 provide the observed monthly and seasonal mean climatic conditions, energy fluxes, GPP, R, and NEE for the study period from January 2002 through January 2006. The TNF averages 1920 mm per year of precipitation with a mean dry season of 5 months duration (months with <100 mm precipitation), typically extending from July 15 to December 15 [Parrotta et al., 1995]. This site is in the 27th percentile (±2–3%) for both annual precipitation and length of the wet season in the Amazon basin [Saleska et al., 2003; auxiliary material]. There is a regional minimum in annual precipitation in the North-Central Amazon, the location of the TNF, because the propagating sea-breeze front that provides an important trigger for convective precipitation arrives at night [Garstang et al., 1994; Mitchell et al., 2003]. There was also a tendency for precipitation to occur in the late afternoon (1300–1500, local time (LT)) during all seasons,
due to convective activity stimulated by surface heating. Climatic conditions during our four years of observation were sufficiently variable to allow us to examine both seasonal and inter-annual variability, but did not include major climatic extremes or significant El Niño events.

Meteorology in the TNF is characterized by persistent trade winds [Lu et al., 2005]. Winds at the top of the tower (64 m) were predominantly from the east and northeast. During the afternoons, a westerly river breeze sometimes developed due to differential heating between the forest and the Tapajo River. The river breeze circulation was strongest during hot dry season afternoons, but was also present during dry afternoons in the rainy season. River breezes on average lasted 1.7 hours and developed during approximately 28% of the days.

The mean daily (24-hour average) wind speed recorded at the top of tower by the sonic anemometer was 2.1 ± 0.01 m s⁻¹ during the wet season and 2.2 ± 0.01 m s⁻¹ during the dry season. The mean daytime (0700–1500, LT) u* was 0.42 ± 0.004 and 0.44 ± 0.006 m s⁻¹ during the wet and dry seasons, respectively. The mean nighttime (2300–0500, LT) u* was 0.21 ± 0.004 m s⁻¹ for both the wet and dry seasons.

Observed net radiation flux (Rn) and temperature were higher during the dry season (Tables 2 and 3, Figures 2 and 3). Latent heat flux (LE) and vapor pressure deficit (VPD) closely followed the diel patterns in Rn and temperature. During the study period, the daily mean temperature was 25.9 ± 0.74 °C, with mean daily minimum and maximum temperatures of 23.5 ± 0.05 °C and 29.0 ± 0.08 °C, respectively. Air temperature did not follow a symmetric diurnal cycle. Heating was rapid after sunrise (0600 local time), with slow cooling in the afternoon after 1300 (LT). The observed mean RH and mixing ratio of water vapor were 78.2 ± 0.1 % and 16.2 ± 0.07 g H₂O kg dry air⁻¹ over the study period, respectively.

### 3.2. Energy Balance

Net radiation flux (Rn) at the surface can be partitioned into ground heat flux (G), changes in biomass and canopy air heat content (S), atmospheric sensible (H) and latent (LE) heat fluxes, and ecosystem respiration (R).

![Figure 2](image-url)
latent heat (LE) fluxes, and net energy exchange due to precipitation inputs (\(\Delta E_p\), see below). Energy balance closure dictates that the sum of LE and H be equivalent to other energy sources and sinks such that

\[
R_n - G - S = LE + H + \Delta E_p.
\]

Energy balance closure is an important criterion used to assess the reliability and accuracy of surface flux measurements. G was not measured at this site, but has been estimated to be of order 3 W m\(^{-2}\) during the daytime (N. Hasler and R. Aviszar, What controls Amazon evapotranspiration?, submitted to Journal of Hydrometeorology, 2007, hereinafter referred to as Hasler and Aviszar, submitted manuscript, 2007) with its 24-hour integral approaching 0. In Amazonian ecosystems, where the quantity of biomass is very large, S has been estimated to be approximately 5–10% of incoming net radiation [Moore and Fisch, 1986], but also averages to 0 over daily intervals. S was estimated for this site using the empirical relationship reported in Moore and Fisch [1986] for a tropical forest near Manaus, Brazil.

To assess energy balance closure, we examined the slope of an orthogonal distance regression of daytime hourly turbulent heat fluxes (LE + H) versus the total available energy (\(R_n - S\)) for all daytime hourly measurements (neglecting ground and \(\Delta E_p\) fluxes); average closure was 85% (\(\pm 0.08\)) using this method. Energy closure was higher during the dry season (88 ± 0.1%) than the wet season (83 ± 0.08%). The seasonal closure difference may be a measure of unquantified heat exchanged by precipitation. For example, a 10 mm hr\(^{-1}\) rain event, with the water ten degrees cooler than ambient air, could result in an apparent loss of 116 W m\(^{-2}\) from the ecosystem that is not captured in this analysis. On an annual basis the energy flux due to rain (\(~2000\) mm yr\(^{-1}\)) is of order 3% of the total net radiation, and will have a larger impact in the wet season. Measurement artifacts such as sensor separation and finite volume averaging also result in small, consistent losses in LE and H fluxes [Finnigan, 2004]. Given the overall consistency between wet and dry energy closure results, there is no reason to suspect our fluxes are significantly biased on seasonal timescales. Our observed 15% lack of closure in hourly data is similar to observations at most flux tower sites; global average closure was found to be 79% [Wilson et al., 2002] and 82% within the Amazon tower sites (Hasler and Aviszar, submitted manuscript, 2007). The slope of the 24-hour energy closure (LE + H vs. Rn) was 93% (\(\pm 2.8\)), using only days with complete data coverage (more common in the wet season).

### 3.3. Ecosystem Carbon Fluxes

The annual (January–December) carbon balances at this site were 2677 ± 488, 906 ± 491, −221 ± 453, and 392 ± 449 kg C ha\(^{-1}\) yr\(^{-1}\), for 2002–2005, respectively, indicating a small net source of carbon to the atmosphere over the period, declining to approximate carbon balance over four
The complete record analyzed here confirms the seasonal patterns initially reported for the TNF by Goulden et al. [2004] and Saleska et al. [2003]. During the wet season R was generally greater than GPP, resulting in a net carbon loss to the atmosphere (Tables 2 and 3, Figures 4 and 5). During the dry season the reverse was more common, with GPP exceeding R resulting in net carbon uptake from the atmosphere.

The mean annual ecosystem respiration was 8.6 ± 0.11 μmol m⁻² s⁻¹, with a mean of 9.2 ± 0.15 and 7.7 ± 0.15 μmol m⁻² s⁻¹ for the wet and dry seasons, respectively. Maximum respiration was observed during the mid-wet season in March and minimum respiration was observed during the late dry season in October (Table 2 and Figure 5). The mean annual GPP was 8.3 ± 0.11 μmol m⁻² s⁻¹ with no statistically significant seasonal difference in carbon uptake, 2614 ± 93 kg C ha⁻¹ month⁻¹ and 2653 ± 79 kg C ha⁻¹ month⁻¹ for the wet and dry seasons, respectively.

The total ecosystem R was lower during the dry season, but the decline in R typically began during the latter part of the wet season, in synchrony with the decline in canopy uptake. R tended to remain low throughout the dry season even as canopy uptake increased. This observation appears to highlight differential responses of the autotrophic and heterotrophic components of R. Autotrophic R can be assumed to increase with increasing GPP. Hence, reduction of R in the dry season is very likely to represent moisture limitations on heterotrophic R. Over four years, the TNF was a source of carbon to the atmosphere with an observed mean loss of 890 ± 220 kg C ha⁻¹ yr⁻¹.

3.4. Ecosystem Water Fluxes

Observed ET ranged widely, from 0.67 to 6.24 mm day⁻¹, with average rates of 2.89 ± 0.15 and 3.41 ± 0.18 mm day⁻¹ for the wet and dry seasons, respectively. The annual mean total was 1135 mm. Across the four measurement years, ET consistently increased at the start of the dry season and remained elevated throughout the entire dry season (Figure 3). ET rates were within the range observed at other Amazonian flux sites (see Amazon-wide comparisons in Hutyra et al. [2005]), but the data were significantly lower than modeled ET reported by Nepstad et al. [2004] and Lee et al. [2005]. The annual fraction of precipitation lost through ET was fairly constant during the study period at 0.53 (1116 mm/2111 mm), 0.64 (1114 mm/1740 mm), 0.49 (1137 mm/2311 mm), 0.51 (1123/2201) for 2002–2005, respectively. The ratios of evaporation to precipitation during the dry seasons of 2002, 2003, 2004, and 2005 were 1.81 (503.3 mm/278.5 mm), 1.16 (521.8 mm/448 mm), 1.28 (514.4 mm/402.4 mm), and 1.40 (535.7 mm/382.9 mm), respectively. Dry season ET was insensitive to dry season precipitation, being nearly constant across years even though dry season precipitation varied by 40%.

There was no statistically significant difference observed between the wet and dry seasons in the slope of LE and H versus Rn (Figure 6). The mean annual evaporative fraction (LE/Rn) was 0.62. This invariance contrasts markedly with data reported by Malhi et al. [1998, 2002], who observed significant seasonal differences in the evaporative fraction in an Amazonian forest near Manaus, Brazil, that actually receives more rainfall and has a shorter dry period. The observed patterns are consistent with the findings of da Rocha et al. [2004] for the nearby tower at km 83.

3.5. Independent Estimates of Carbon Flux

It is critical to independently validate carbon flux measurements in order to ensure accurate cumulative sums and to examine the mechanisms controlling exchange. Biases in day/night measurements of CO₂ flux could
significantly affect estimates of the overall carbon balance. A potential source of bias is the prevalence of weak vertical mixing during the nighttime hours, leading to a violation of the assumption of horizontal homogeneity required for eddy flux measurements and to ‘lost flux’ associated with horizontal advection [Finnigan, 2004]. We used three independent approaches to ensure unbiased data for nighttime fluxes and to validate flux measurements: (1) filtering the data according to $u^*$ values to correct for underestimation of nighttime fluxes; (2) analysis of annual and seasonal light response relationships between PAR and NEE to derive independent estimates of nighttime NEE, avoiding any use of nighttime data or $u^*$ filtering; (3) estimation of nighttime NEE by similarity of CO$_2$ with $^{222}$Rn.

Figure 5. Monthly time series (49 months) of (a) net ecosystem exchange of CO$_2$ (NEE, $\mu$mol m$^{-2}$ s$^{-1}$), (b) ecosystem respiration (R, $\mu$mol m$^{-2}$ s$^{-1}$), and (c) gross primary production (GPP, $\mu$mol m$^{-2}$ s$^{-1}$). The shading patterns within the bars indicate the season.

Respiration is a biological process that should be largely independent of the turbulence intensity. Since measured NEE decreased in calm conditions (Figure S2), there appears to be some lost flux. Approximately 57% of the nighttime hours at this site were calm, with $u^* < 0.22$ m s$^{-1}$. We corrected for lost flux by filtering calm night periods and replacing the data with the mean value for proximate well-mixed time periods (defined as $u^* \geq 0.22$ m s$^{-1}$, see Saleska et al. [2003] and Hutyrà et al. (submitted manuscript, 2007) for further discussion of $u^*$ corrections and the relationship between canopy CO$_2$ storage and turbulence). Note that the prevalence of strong turbulence (high $u^*$) in both daytime and nighttime is higher at the TNF than observed at many Amazonian flux towers, giving better mixing and fewer gaps in the nighttime flux [cf. Kruif et al., 2004]. The observed mean nighttime NEE with $u^*$ filtering was $9.2 \pm 0.15$ and $7.7 \pm 0.15$ $\mu$mol m$^{-2}$ s$^{-1}$ for the wet and dry seasons, respectively; were no $u^*$ filter applied the respective mean nighttime NEE would be $7.1 \pm 0.09$ and $5.8 \pm 0.10$ $\mu$mol m$^{-2}$ s$^{-1}$.

We examined NEE-light relationships (Figure 7) using a nonlinear least squares approximation (hyperbolic function)

$$NEE = a_1 + \frac{a_2 \times PAR}{a_3 + PAR}$$

fitted to NEE and PAR. We excluded data for PAR $\leq 40$ $\mu$mol m$^{-2}$ s$^{-1}$, since these data points often correspond to periods of low turbulence and rapidly changing light levels, resulting in large uncertainties. The intercept, $a_1$, of this fit provides an independent estimate of mean ecosystem R (limit of equation (5) as PAR $\rightarrow 0$). The annual mean value
of a1 was 8.9 ± 0.6 μmol m⁻² s⁻¹, based on all available data (no u* filter applied), and statistically indistinguishable from the mean nighttime u* filtered NEE (8.6 ± 0.13 μmol m⁻² s⁻¹). Note that the respective data sets are completely disjoint. Seasonal comparisons between a1 and u*-filtered mean nighttime NEE also agreed within 5% (Figure S3).

3. Data for ²²²Rn can potentially define rates of forest-atmosphere exchange, since ²²²Rn is conserved after emission from the soil (apart from slow radioactive decay). Martens et al. [2004] independently assessed raw and u* corrected eddy flux NEE measurements at night by comparing CO₂ eddy flux data with CO₂ fluxes inferred from ²²²Rn profiles and ²²²Rn soil flux measurements. Nighttime NEE derived from ²²²Rn was found to be 9.0 ± 0.99 μmol m⁻² s⁻¹ for the wet season (June–July 2001) and 6.4 ± 0.59 μmol m⁻² s⁻¹ in the dry season (November–December 2001), agreeing very well with u* filtered NEE measurements during the same period (8.65 ± 1.07 and 6.56 ± 0.73, respectively) [Martens et al., 2004].

The independent light-curve and ²²²Rn based estimates of nighttime NEE both agree extremely well with the u*-filtered nighttime flux measurements. Failing to apply a u* filter to the data would have changed the annual sum of carbon exchange from a small carbon source to a significant carbon sink, almost 10 Mg C ha⁻¹ yr⁻¹. This value would also markedly disagree with bottom-up estimates for this site [Rice et al., 2004; Saleska et al., 2003; Hutyra et al., submitted manuscript, 2007].

4. Discussion

4.1. Controls on NEE

The carbon balance of an ecosystem is the result of disturbance and recovery dynamics over timescales of years...
and decades [Saleska et al., 2003; Rice et al., 2004; Vieira et al., 2004], upon which is superimposed the influence of weather anomalies on seasonal and annual timescales. Figure 4 shows the cumulative annual cycles of NEE, highlighting the dominance of ecosystem respiration throughout the early portion of the calendar year (wet season) as the forest lost carbon to the atmosphere. By September, increases in canopy uptake generally began to dominate and the forest turned into a carbon sink for the rest of the dry season. The transition back to a net carbon source followed the arrival of wetter weather.

Climate anomalies exerted strong control on the inter-annual variations in net carbon balance. In 2005, carbon losses in the wet season were relatively small and the transition to carbon uptake was very abrupt, and by November the year was on track to be a significant carbon sink. But, the early arrival of the wet season, with significant November and December rainfall, reversed the carbon uptake and the site was instead an overall carbon source in 2005 (Figures 4 and 5). In January 2003, low precipitation, totaling only 27 mm, resulted in reduced respiration rates. However, GPP rates remained high, leading to a carbon sink for the month despite the seasonal norm (Figures 4 and 5). The greatest variability in monthly total NEE was observed during the late dry season and early wet season (November–January). Respiration rates were the most variable and sensitive to precipitation and temperature anomalies (Figure 5).

4.2. Controls on Gross Primary Production

Many process-based biogeochemical models [e.g., Botta et al., 2002; Tian et al., 1998] predict that moisture limitation during the dry season should provide a strong constraint on canopy carbon uptake in tropical forests like the TNF. Four years of observations at the TNF do not support this paradigm. Uptake was indeed reduced early in the dry season, but the decline began before the onset of the dry weather. Moreover, uptake started to increase in the driest period, well before the onset of the rainy season (Table 2; Figures 4 and 5). The forest maintained high rates of photosynthesis throughout the year because of adequate water supplies, high year-round temperatures, and high light levels. Goulden et al. [2004] observed a similar seasonal pattern in photosynthesis at a nearby forest site between July 2000 and July 2001.

Peak litterfall rates were observed at the TNF in August [Rice et al., 2004], early in the dry season, and the flush of new leaves across the Basin also occurred in the dry season, August–October (Figure 8) [Huete et al., 2006; Rivera et al., 2002]. Younger leaves have higher photosynthetic efficiency [Freeland, 1952] and hence it is not
surprising that higher rates of GPP were observed in the months following leaf-out in the dry season. Previous work by Wright and van Schaik [1994] also showed that tropical plants produce new leaves when irradiance is maximized. 

To quantify the phenology effects on GPP at this site, we calculated “canopy photosynthetic capacity” ($P_c$) as the mean monthly GPP in a fixed light interval (PAR 725–875 μmol m$^{-2}$ s$^{-1}$), and compared the time series of this quantity with leaf litterfall rates and with remotely sensed vegetation greenness (enhanced vegetation index, EVI) at the TNF [Huete et al., 2006]. We examined $P_c$ to remove the influence of seasonal differences in incoming radiation. Figure 8 shows that leaf litterfall rates were strongly correlated with $P_c$ ($r^2 = 0.76$ or 0.83, for lags of 0 or 1 months, respectively). In contrast, EVI correlated weakly with $P_c$, explaining at best only 56% of the observed variance with a long lag (3 months). EVI, lagged by 2 months, was somewhat better correlated with monthly litterfall ($r^2 = 0.63$). The temporally lagged correlations in EVI and/or litterfall are not surprising since it takes time for the leaves to fully elongate and develop their pigmentation. Note that total GPP, across all light levels, also correlated with litterfall ($r^2 = 0.63$, lagged by 2 months) and EVI ($r^2 = 0.40$, leading by 1 month).

During the late dry season there are also increased aerosol loadings due to land clearing and agricultural activities, resulting in higher diffuse light levels. P. H. F. Oliveira et al. (The effects of biomass burning aerosols and clouds on the CO$_2$ flux in Amazonia, submitted to Tellus, Ser. B, 2007, hereinafter referred to as Oliveira et al., submitted manuscript, 2007) observed maximum aerosol loading at the TNF between September and November. Higher photosynthetic rates have been observed under diffuse light conditions [Gu et al., 2003; Oliveira et al., submitted manuscript, 2007]. Either or both leaf replacement and aerosol light scattering may account for increased $P_c$ in the late dry season (October–December, Figure 7). There was significant inter-annual variation in both EVI and $P_c$, see Figure S4 for the full available time series.

### 4.3. Controls on Ecosystem Respiration

Ecosystem respiration is the sum of CO$_2$ released by plant leaves, stems, and roots (autotrophic respiration), and CO$_2$ released through decomposition of organic material (heterotrophic respiration). Temperature and moisture are key environmental factors regulating respiration rates, but the interaction among these parameters is still poorly understood [Raich and Schlesinger, 1992; Trumbore, 2006]. Temperature and soil moisture are typically inversely correlated, but both factors simultaneously influence R by affecting enzyme activity, diffusion of solutes and O$_2$, growth of root tissue, and microbial populations [Davidson et al., 2006]. Eddy covariance data cannot distinguish the components of respiration. However, our long data set from the TNF does allow us to examine the aggregate effects of climatic variability on total ecosystem respiration, over timescales from hourly to inter-annual.

Relationships between R and temperature have been reported in many ecosystems, and ecosystem models often use exponential relationships to describe these data, with $Q_{10}$ values typically between 1 and 2 [e.g., Davidson et al., 2006]. But decomposition of organic material in tropical forest soils is known to have a relatively low temperature sensitivity [Davidson and Janssens, 2006]. At the TNF, there was no statistically significant relationship between nighttime CO$_2$ flux and ground or canopy temperature, or precipitation, over any time interval from hourly to weekly (Table 4). Davidson et al. [2004], working at nearby site (~5 km), also found no significant relationship between soil volumetric water content and observed soil CO$_2$ respiration rates using chamber methods. The absence of a significant relationship between temperature and ecosystem R could be an artifact of high mean temperature, with canopy and ground mean temperatures averaging 24.8°C and 24.6°C, respectively, or of the small temperature range seasonally, diurnally, and during the nighttime. It also possible that the entire temperature range is within a broad optimum for this ecosystem or that the temperature responses of multiple processes may cancel when aggregated to the ecosystem scale. The observations imply that models of tropical forests which include an exponential relationship between respiration and temperature may over-predict the temperature sensitivity of respiration rates at the ecosystem level.

When averaged on longer timescales, temperature and precipitation were significant correlates of total ecosystem respiration and a temperature regression could indeed explain the most significant portion of the total observed variance (Table 4). But, respiration was negatively correlated with temperature and positively correlated with precipitation, and the apparent relationship between $R$ and temperature arises because temperature and precipitation are negatively correlated. We examined the intercept values ($a_0$) of morning versus afternoon light-curve extrapolations (equation (5)) and found no significant difference in the respiration estimates in the dry season, although temperature differences were near their maximum (Figure S5). In contrast, during the wet season we found that morning respiration estimates were higher than the afternoon estimates in three of four observed wet seasons (Figure S5). Higher morning respiration highlights the dominance of
moisture in controlling heterotrophic respirations rates since nighttime precipitation is very common while morning temperatures were lower. We conclude that the negative respiration-temperature correlation is likely a simple artifact that arises because wet seasons, which have higher respiration rates, are cooler than dry seasons.

Maximum litterfall rates (leaves, twigs, and fruits) were observed shortly after the onset of the dry season in August and September (Figure 8) [Rice et al., 2004]. Tropical forest litter typically has a short turnover time (less than 1 year [Brown and Lugo, 1982]), but during the dry season, following the peak input of litter, moisture levels are low in soil and litter. Hence, ecosystem respiration rates remain low, even though substrate abundance, temperatures, and canopy metabolic rates were highest in the dry season.

Chambers et al. [2004] estimated the mean ecosystem respiration rate to be 7.8 \( \mu mol \ m^{-2} \ s^{-1} \) at a site near Manaus, Brazil by measuring individual components of ecosystem respiration, compared to 8.4 \( \mu mol \ m^{-2} \ s^{-1} \) (7.5–9.4 95% CI) using eddy covariance method at that site. Using chamber-based methods, Chambers et al. [2004] estimated a mean soil respiration rate of 3.2 \( \mu mol \ m^{-2} \ s^{-1} \) at the Manaus site and reported that both soil respiration and total ecosystem respiration declined with increasing soil moisture, opposite to our observations. Chambers et al. [2004] speculated inadequate oxygen supplies in saturated soils led to lower respiration rates. Soil respiration measured at the TNF (R. K. Varner et al., manuscript in preparation, 2007) averaged 2.63 \( \mu mol \ m^{-2} \ s^{-1} \) annually and 2.91 and 2.29 \( \mu mol \ m^{-2} \ s^{-1} \) for the wet and dry seasons, respectively, and showed an increase with precipitation and a negative correlation with temperature, in harmony with our data for total ecosystem R. Thus, R at these two sites showed opposite seasonality, peaking during the wet season at the TNF but in the dry season at Manaus.

It is not known why these sites exhibit different seasonality in respiration. The TNF has much more coarse woody debris (CWD; 48 ± 5.2 Mg C ha\(^{-1}\) [Rice et al., 2004; G. W. Santoni et al., manuscript in preparation, 2007]) than in Manaus (10.5 Mg C ha\(^{-1}\) [Chambers et al., 2000]; 15.7 ± 4.1 Mg C ha\(^{-1}\) [Nascimento and Laurance, 2004], assuming that 1 kg dry biomass = 0.5 kg C biomass). In the TNF, CWD respiration was estimated to be a very significant component of the overall respiration budget contributing 1.2 ± 0.3 \( \mu mol \ m^{-2} \ s^{-1} \) (see Hutyra et al. submitted manuscript, 2007) for a full breakdown of the TNF respiration budget. In contrast, CWD respiration estimates from Manaus are significantly smaller, contributing only ~0.50 \( \mu mol \ m^{-2} \ s^{-1} \) [Chambers et al., 2004]. Seasonal patterns of CWD respiration are very poorly quantified for the tropics, but it is possible that the moisture and temperature responses of CWD respiration could differ significantly from soil R. The combination of a longer dry season and a larger stock of CWD at the TNF may contribute to changes in CWD respiration and help explain the seasonal differences versus Manaus. It is also possible that a different moisture optimum exists in Manaus due to the shorter dry season and greater annual rainfall. Further, topographic and soil differences between the Manaus and TNF sites are likely to also contribute to the opposite seasonal respiration patterns. The Manaus study site is located within an area of undulating topography with often inundated soils in the low-lying areas [Araújo et al., 2002] while the TNF has very little topographic variation, no soil inundation, and an extremely deep water table. The physical reason for the seasonal differences in respiration remains an open question in need of further research.

### 4.4. Is Forest Growth Water Limited?

Seasonal water limitations have the potential to reduce forest growth and place the forest at risk for fire. Future climate scenarios suggest that temperatures in the Amazon may increase while precipitation decreases [Fung et al., 2005], likely decreasing water availability and increasing drought and flammability. To assess the current sensitivity of this forest to water limitations, we looked at the patterns in water flux, seasonal evaporative balances, water-use efficiencies, and light-use efficiency.

ET rates consistently increased at the start of the dry season and remained elevated throughout the period of maximum radiation inputs (Figures 2 and 3). Water losses consistently exceeded inputs during the dry season, large stores of water are evidently accessible to the trees. In the case of 2002, approximately 225 mm of water was withdrawn from storage during the dry season. If we adapt as representative the plant available water profile measured in a similar soil by Nepstad et al. [1994], the forest had to extract water from depths in excess of 4 m to support the observed dry season ET rates. The higher ET rates and the nearly inelastic total ET in the dry season are both strong indicators of adequate water availability at the TNF with the current climate.

Ecosystem water-use efficiency (WUE) can be defined as the ratio of GPP to \( F_{H2O} \) (carbon uptake/water loss). Elevated values of WUE could indicate water stress as the scarce resource (water) is conserved. But, the mean observed WUE was 4.5 and 3.7 \( \mu mol \ CO_2/mmol \ H_2O \) for the wet and dry season, respectively, showing the opposite trend. Although this result is consistent with this ecosystem not experiencing seasonal water stress, it must be interpreted cautiously. Changes in the WUE can result from a change in either the canopy carbon uptake or \( F_{H2O} \). As the dry season approached, \( F_{H2O} \) and the vapor pressure deficit started to increase while the GPP started to decrease, resulting in a lower overall dry season WUE. The WUE started to increase again in October when canopy carbon uptake increased, while the \( F_{H2O} \) remained high (Tables 2 and 3).

Both light-use efficiency (LUE) and WUE were significantly higher in the morning than afternoon (Figure 9). The diel patterns in the LUE and WUE are consistent with afternoon GPP being inhibited. The standard paradigm is that as VPD increases, plant water stress will increase and stomatal conductance will decrease, resulting in higher WUE and lower LE and LUE. However, at this site the LE remains high in the afternoon (Figure 2) and the overall evaporative fraction increased along with VPD, likely indicating abundant water supplies. The fraction of water lost through transpiration may change diurnally, but the LE measurements can not be readily separated into the component processes. It is possible that the apparent afternoon reductions in GPP were due to differences in autotrophic respiration rates, but the analysis of light-curve intercepts (Figure S5) does not support that interpretation. The morning and afternoon differences in WUE and LUE are more
likely due to limitation on stem conductance, plant circadian rhythms [Doughty et al., 2006], metabolic cycles (e.g., respiration associated with sugar transport), or enzymatic limitations.

5. Summary and Conclusions

[55] In this study we critically assessed flux measurements of CO$_2$ and H$_2$O and examined energy closure to ensure the validity of the observations, then we examined the controls on carbon and water exchange in an evergreen tropical rain forest. We found no significant signs of water limitation on photosynthesis: trees had adequate water supplies throughout the 5 month dry season. ET responded strongly to radiative drivers year-round and was insensitive to dry season precipitation totals. Observed dry season evaporative losses significantly exceeded precipitation inputs, drawing up to 225 mm of water from water reserves that had to extend many meters in depth. Evidently the annual input of precipitation and the capacity of the plants to use stored water over considerable depth provide the key to maintaining this closed canopy equatorial forest despite long periods of low rainfall.

[56] We found that the seasonal course of canopy photosynthesis was largely controlled by phenology and light. Canopy photosynthetic efficiency declined before leaf senescence (late wet season) and increased after new leaf elongation (mid-dry season). Unfortunately, the EVI parameter did not capture this pattern. Phenological control of the timing of peak carbon uptake capacity ($P_c$) again highlights adequate water availability and suggests that the assemblage of trees in this forest may have been selected to optimize for light, not water. The dominant influence of phenology versus water stress is a significant surprise for this forest.

[57] Climate anomalies exerted a strong influence on net carbon exchange, principally through effects on R. Ecosystem R was lower during the dry season due to moisture limitations on heterotrophic respiration as evidenced by enhanced dry season GPP and $P_c$ rates. We did not find a significant relationship between temperature and R on short timescales. The lack of temperature dependence raises uncertainty about the appropriateness of using Q-10 type relationships in ecosystem models of tropical rain forests. The largest variations in R, photosynthesis, and net carbon exchange were observed during the dry-to-wet season transition.

[58] This forest currently does not exhibit signs of water limitations, with enough water to satisfy growth requirements. It was a small overall carbon source to the atmosphere, with the efflux rate declining over the period of study, consistent with the long-term ecosystem disturbance and recovery dynamics as proposed by Rice et al. [2004] and with the large contribution of CWD respiration to the observed high rates of respiratory carbon losses. Live biomass stocks have increased significantly over the study period while CWD stocks have decreased (G. W. Santoni et al., manuscript in preparation, 2007).

[59] If precipitation rates were to decrease by a small amount, but water supplies remained adequate for the trees, it is possible that the net carbon uptake could increase due to higher insolation and slower heterotrophic respiration. However, a reduction in decomposition from drier conditions could result in increased flammability due to a build up of fuel. Alternatively, if the amount of available water for the trees were to decrease through logging (causing soil compaction), higher temperatures (increasing the evaporative demands), or large decreases in precipitation (slowing recharge of deep water reservoirs) the flammability of this forest might increase and the forest may convert to a fire adapted vegetation type. Accurate predictions of future climate and land-use changes require capturing these critical dependencies on precipitation and on ecosystem structure and function.
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