EMILY ANE DIONIZIO DA SILVA

INFLUENCE OF CLIMATE, FIRE AND PHOSPHORUS IN THE DYNAMICS OF VEGETATION IN THE AMAZON-CERRADO BORDER SIMULATED WITH INLAND MODEL

Dissertação apresentada à Universidade Federal de Viçosa, como parte das exigências do programa de Pós-Graduação em Meteorologia Aplicada, para obtenção do título de *Magister Scientiae*.

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EMILY ANE DIONIZIO DA SILVA

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BIOGRAFIA

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С	Carbon
cm	Centimeter
m	Meters
$m^2 m^{-2}$	Square meter per square meter
mm yr ⁻¹	Millimeter per year
mg	Milligram
mg L ⁻¹	Milligram per liter
mg dm ⁻³	Milligram per cubic decimeter
mg kg ⁻¹	Milligram per kilogram
μmol	Micromolar
m²	Square meter
S	Seconds
Р	Phosphorus
Ν	Nitrogen
kg	Kilogram
kg m ⁻²	Kilogram per square meter
yr	Year

LIST OF SYMBOLS

LIST OF FIGURES

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LIST OF ACRONYSMS

С	Clay
BESM	Brazilian Earth System Model
CA	Average climate (climatology 1961-1990)
CENTURY	Model of plant-soil nutrient cycling
CLM	The Community Land Model
CMAP	Maximum capacity of the soil adsorption
CO_2	Carbon Dioxide
CTEM	Canadian Terrestrial Ecosystem Model
CV	Variable climate (database 1948-2008)
DGVMs	Dynamic Global Vegetation Models
F	Fire
GPP	Gross primary productivity
HCL	Hydrochloric acid
IBIS	Integrated BIosphere Simulator
IGBP	International Global Biosphere Programme
INLAND	Integrated Model of Surface Process
ISAM	Integrated Science Assessment Model
JPL	Lund-Potsdam-Jena Dynamic Global Vegetation Model
LAIupper	Upper canopy leaf area index
LAI _{lower}	Lower canopy leaf area index
LAI _{dry}	Dry season leaf area index (MODIS)
LSM	Land Surface Model
MODIS	Moderate Resolution Imaging Spectroradiometer
NPP	Net primary production
P-Mehlich-1	Phosphorus extracted by Mehlich-1 extractor
P_m1_est	Estimated Phosphorus extracted by Mehlich-1 extractor
P _{Adc}	Phosphorus added in each soil sample
PC	Control Phosphorus
PFT	Plant functional type
PG	Global Phosphorus
PR	Regional phosphorus
Prem	Remanescent phosphorus

Presin	Phosphorus extracted by Resin
P _{total}	Phosphorus total in the soil
Rubisco	Ribulose-bisfosfato carboxilase oxigenase
UNEMAT	State University of Mato Grosso
V _{max}	Maximum rate of carboxylation Rubisco
PROBIO	Brazil - National Biodiversity (PROBIO) Project

ABSTRACT

SILVA, Emily Ane Dionizio da, M. Sc. Universidade Federal de Viçosa, February, 2015. Influence of climate, fire and phosphorus in the dynamics of vegetation in the Amazon-Cerrado border simulated with INLAND model. Adviser: Marcos Heil Costa. Co-Adviser: Júlio Cesar Lima Neves.

Studies to assess the main factors responsible for the dynamics of vegetation in the Amazon such as climate and / or deforestation suggest the possibility of the Amazon rainforest may not resist a potencial change in rainfall patterns, and be converted to an ecosystem of more sparse vegetation like savanna, which gave rise to the term "Amazon Savannization". Some of these initial studies, however, used only future climate scenarios, neglecting in their majority the fire effects and especially the effect of nutrient limitation on vegetation dynamics. Recently, in the context of climate modeling, advancements have been made in an attempt to build models that incorporate the nutritional restriction effect and the fire to assess and measure their effects on vegetation at a large scale. In this work, the main objective was to evaluate the isolated and combined effects of climate (inter annual variability and CO₂), fire (F) and nutritional limitation by phosphorus (P) using the dynamic vegetation model INLAND in the transition between forest-Cerrado. The INLAND model is the first Brazilian model to simulate the vegetation dynamics incorporating these factors. The INLAND characterizes the vegetation through plant functional types, which combined differently give rise to biomes. The nutritional limitation is established through a linear function of the total phosphorus content in the soil and the maximum carboxylation capacity of Rubisco enzyme (V_{max}). The Rubisco is the main enzyme responsible for regulating carboxylation rates in plants with the C3 photosynthetic mechanism type acting directly on net primary productivity (NPP) and the increase of biomass. Two regional the datasets of V_{max} to be used as input data of the simulations, based on regional (PR) (field measurements) and global data (PG) of total P content in the soil. In the design of numerical experiment, 24 simulations were performed for different combinations of climate factors, P, F and CO₂ were obtained. To validate the simulations, regional datasets of dry season leaf area index (LAI), and biomass were compared against simulated values by the INLAND model. The results showed that isolation, the effect of fire on the NPP in the Cerrado is positive and higher than the effects of nutritional restriction, contrary to what happens in the Amazon biome where the P acts to reduce the NPP. The CO₂ when evaluated in combination with the effects of fire acted as an

attenuator increasing the NPP in the transition region. With respect to the climate, the use of inter-annual variability improved representation of biomass and LAI simulated by INLAND. The LAI and biomass simulated values showed good correlation with Amazon-Cerrado observed gradient. The best representation of the vegetation composition by INLAND was found when the effects of $PG+F+CO_2$ were combined.

RESUMO

SILVA, Emily Ane Dionizio da, M. Sc. Universidade Federal de Viçosa, Fevereiro de 2015. Influência do clima, fogo e fósforo na dinâmica da vegetação na fronteira Amazônia-Cerrado simulada pelo modelo INLAND. Orientador: Marcos Heil Costa. Coorientador: Júlio Cesar Lima Neves

Estudos realizados para avaliar os principais fatores responsáveis pela dinâmica da vegetação na Amazônia tais como clima e/ou desmatamento sugerem a possibilidade da floresta Amazônica não resistir a uma eventual mudança do regime de chuvas, e ser transformada em um ecossistema de vegetação mais esparsa, do tipo savana, o que deu origem ao termo "Savanização da Amazônia". Parte destes estudos iniciais no entanto utilizavam apenas cenários de climas futuros, negligenciando na maior parte das vezes o efeito do fogo, e principalmente o efeito da limitação nutricional na dinâmica da vegetação. Recentemente, no âmbito da modelagem climática, grandes avanços têm sido realizados na tentativa de se construir modelos que incorporem o efeito da limitação nutricional e do fogo de forma que se possa avaliar e dimensionar os efeitos dos mesmos na vegetação em larga escala. Neste trabalho, o principal objetivo foi avaliar os efeitos isolados e combinados do clima (variabilidade interanual e CO_2), fogo (F) e limitação nutricional por fósforo (P) utilizando o modelo de vegetação dinâmica INLAND na transição entre Amazônia-Cerrado. O modelo INLAND é o primeiro modelo brasileiro que permite simular a dinâmica da vegetação incorporando estes fatores. O INLAND caracteriza a vegetação através de tipos funcionais de planta, que combinados de forma diferente dão origem aos biomas. A limitação nutricional no modelo é estabelecida através de uma função linear entre o teor de fósforo total no solo e a capacidade máxima de carboxilação da enzima Rubisco (V_{max}). A Rubisco é a principal enzima responsável por regular as taxas de carboxilação nas plantas com mecanismo fotossintético do tipo C3 atuando diretamente na produtividade primária líquida (NPP) e no incremento de biomassa. Dois mapas de Vmáx foram elaborados para serem utilizados como dados de entrada nas simulações, baseados em dados regionais (PR) (medidas de campo) e globais (PG) de teor de P total no solo. No delineamento do experimento numérico, 24 simulações foram realizadas de forma que diferentes combinações entre os fatores clima, P, F e CO2 fossem obtidas. Para validação das simulações, um mapa de índice de área foliar (LAI) MODIS e dados de biomassa foram contrastados com valores simulados pelo modelo INLAND. Os resultados demonstraram que isoladamente, o efeito do fogo sobre a NPP no Cerrado é positivo e maior em relação aos efeitos da limitação nutricional, contrariamente ao que acontece na região Amazônica onde o P atua na diminuição da NPP. O CO₂ por sua vez quando avaliado combinadamente aos efeitos do fogo demonstrou-se como um amenizador, aumentando a NPP na região de transição. Com relação ao clima, o uso de uma variabilidade inter-anual melhorou a representação dos valores de NPP, biomassa e LAI pelo INLAND. Os valores de LAI e biomassa simulados apresentaram boa correlação com os dados observados de forma que o gradiente de vegetação Amazônia - Cerrado conseguiu ser representado. A melhor representação da composição da vegetação pelo modelo INLAND foi quando considerado os efeitos de PG+F+CO₂.

GENERAL INTRODUCTION

On a global scale, the distribution of different forms of existing vegetation is in general controlled by the climate where the relations between precipitation and temperature determine the structural patterns of vegetation (Whittaker, 1975). At a regional level however, these structural patterns of vegetation can still depend on local environmental conditions imposed by local variations in soils, and topography.

The boundaries between the Amazon forest and the Brazilian Cerrado, for example, are not only characterized by a seasonal rainfall gradient, but also influenced in large part by the occurrence of fire, availability of water and nutrients in the soil, herbivory and soil characteristics (Lehmann et al., 2011; Murphy and Bowman, 2012; Sankaran et al., 2004). Due to the impossibility of manipulating some of these factors in their natural environments, it is difficult to assess and quantify the degree of interaction between these various factors and to infer how each of them works on the regional vegetation. Therefore, Dynamic Global Vegetation Models (DGVMs) have been used to isolate and manipulate factors like climate, fire and nutrients, to understand the largescale effects of these factors on vegetation (Favier et al., 2004; Hirota et al., 2010; House et al., 2003).

The first modeling studies to assess the function of the dynamics of vegetation in the Amazon were developed solely on the basis of climate, deforestation, and CO₂ fertilization (Betts et al., 2004; Cox et al., 2000; Oyama and Nobre, 2003; Shukla et al., 1990). These modeling studies indicated that the Amazon rainforest might experience eventual changes in rainfall patterns and either transforms in an ecosystem with more sparse, vegetation savanna like, which gave rise to the term "savannization of the Amazon" (Betts et al., 2004; Cox et al., 2000; Oyama and Nobre, 2003; Salazar et al., 2007) or tends towards a seasonal forest (Malhi et al., 2009). These pioneering studies were of paramount importance to the advancement of terrestrial biosphere modeling, but in general, the models commonly used by the scientific community have more realistic representations of temperate and cold climate ecosystems, rather than tropical ecosystems. In tropical regions, the DGVMs neglected important factors such as the effects of fire and nutritional limitations.

In the tropics, fire performs an important ecological role on the Earth's surface and influences the productivity of ecosystems, biogeochemical cycles, and the distribution of biomes by changing the biological activity of plants, including their phenology and physiology (Wright, 2005). Recently, Couto-Santos et al. (2014) demonstrated the effects of the fire occurrence on climate dominance in the dynamics of forest-savanna boundaries, in the northern region of the Brazilian Amazon. These authors showed that, in wet years, the forest advanced over the savannas, while in years with lower rainfall, the forest receded and the savanna expanded due the trend of increased frequency of climatic events, such as drought and fire. In the boundary regions between forest and savanna, fire and climate directly influence the growth rates of the vegetation and, depending on the frequency and intensity, may result in increased mortality of trees in a forest and transformation of an undisturbed forest in an inflammable one (Hoffmann et al., 2012). However, to evaluate the dynamic forestsavanna more broadly, in addition to considering the effects of fire, one also must recognize the importance of nutritional limitations on the growth (productivity) of trees.

Studies support that in tropical forests like the Brazilian Amazon, phosphorus (P) is the main limiting nutrient in the productivity of trees instead of nitrogen (Aragão et al., 2009; Davidson et al., 2004; Goll et al., 2012; McGrath et al., 2001; Mercado et

al., 2009; Paoli et al., 2008; Reich et al., 1995; Silver et al., 1994; Wang et al., 2010). Phosphorus is a nutrient that has not been incorporated in most of DGVMs such as the LPJ (Lund-Potsdam-Jena Dynamic Global Vegetation Model) (Sitch et al., 2003; Smith et al., 2001), the CLM (The Community Land Model) (Oleson et al., 2004), or the ISAM (The Science Integrated Model Assessment) (Jain and Yang, 2005; Jain et al., 1996, 2006). In general, only the nitrogen cycle has been used by DGVMs to limit the productivity in ecosystems. The lack of information on the spatial distribution of phosphorus in the soil in its different forms was a major factor that made the implementation of P nutrient in global climate models difficult (Yang and Post, 2011; Yang et al., 2013). However, recent works such as Quesada et al. (2009, 2010, 2011), Yang and Post (2011) and Yang et al. (2013) provided a comprehensive database of physical and chemical properties of the soil in the Amazon region, and global maps of P content in the soil in its different forms. Then, it is currently possible to implement the P content in the soil and to include the nutrient limitation in the productivity in the DGVM IBIS (Integrated Biosphere Simulator) model (Castanho et al., 2013; Yang et al., 2014).

The IBIS model was used as base for the development of the INLAND (*Integrated Model of Land Surface Processes*). The INLAND model is a Brazilian DGVM able to represent characteristics and important process in South America biomes. The nutritional restriction (Castanho et al., 2013) and fire (Arora and Boer, 2005) on vegetation were recently implemented in this model, which makes it a potential tool for investigating the dynamics of vegetation in Brazilian ecosystems.

Using the INLAND model, this study aimed to evaluate the isolated and combined effects of phosphorus, climate and fire on productivity, leaf area index (LAI),

and biomass increment of vegetation in the Amazon-Cerrado transition, and checking the performance of the model in simulating the Amazon-Cerrado current position. These work is organized in two chapters: Chapter 1: Estimates of total phosphorus in the soil for use in INLAND; Chapter 2: Description and analysis of simulations to evaluate the current position of the Amazon-Cerrado border considering the different phosphorus effects, fire in the current climate scenarios and comparing with leaf area index (LAI MODIS) and biomass values for longitudinal transects in the Amazon-Cerrado boundaries.

CHAPTER 1 – Phosphorus in the soil and plants – development of a phosphorus regional dataset

1.1 INTRODUCTION

In tropical forests like in the Brazilian Amazon, phosphorus (P) is the main limiting nutrient in the productivity of trees instead of nitrogen (Aragão et al., 2009; Davidson et al., 2004; Goll et al., 2012; McGrath et al., 2001; Paoli et al., 2008; Reich et al., 1995; Silver et al., 1994; Wang et al., 2010). In plants, P has an essential role as part of energy-rich compounds such as adenosine triphosphate (ATP) used by most physiological processes such as photosynthesis (Malavolta, 1985). In contrast to nitrogen, which is derived primarily from the atmosphere, phosphorus arises out of rocks and generally presents a high content in younger soils, becoming scarcer in older soils due to weathering processes that lead to the formation of aluminum and iron oxides which are strong adsorbers of P in the soil (Uehara and Gilman, 1981).

This high adsorption capacity is the reason why the tropical soils are strictly poor in available P forms, maintenance P by plants occurring mainly via an intracycling system, where the plant absorbs most of the nutrients from the decomposition of its own litter, stems, trunks and the death of fine roots (Waring and Schlesinger, 1985). Besides the intra-cycling system, a physiological mechanism known as biogeochemical cycling, assists in maintaining the phosphorus content in most plants by carrying P from older, photosynthetically inactive tissues to young growing tissues with high capacity (Poggiani, 1981) as a way to minimize losses to soil P through the adsorption process. According to Switzer and Nelson (1973) about 40% of the N, P and K can be recycled through this mechanism. Recently works such as Mercado et al. (2011) and Quesada et al. (2012) relate the P content in the leaves with the maximum carboxylation capacity of Rubisco enzyme (V_{max}) and the P content in the soil with the woody structure of tropical forests, reinforcing the close relationship between tropical vegetation and the different fractions of P (foliar and soil). In the soil, phosphorus is present in diverse forms that vary according to the chemical nature of the compounds to which it is connected, as well as its binding energy. The different forms of phosphorus found in soil include: labile inorganic phosphorus and organic phosphorus (which are the forms available to plants), occluded phosphorus, phosphorus mineral secondary, phosphorus in the form of apatite and the total phosphorus (P_{total}) (Yang et al., 2013). According to Quesada et al. (2012), total soil P was a better predictor of wood production rates in Amazonia than any of the fractionated organic phosphorus or inorganic phosphorus pools, suggesting that it is not only the immediately available P forms, but probably the entire soil phosphorus pool that is interacting with forest growth on longer timescales.

Based on this finding Castanho et al. (2013) obtained a linear relationship between the P_{total} content in the soil and the V_{max} values for woody plants in the Amazon, so that the amount of net primary productivity was spatially limited by P_{total} content in the soil throughout the Amazon Basin. The nutritional limitation of P in woody plants was restricted to Amazonia due to the lack of P_{total} content measurements for Cerrado.

This chapter has three objectives: (1) to develop a model for estimating the amount of P_{total} from measurements of soil available P content in the Cerrado region on the Amazon-Cerrado transition area located in the state of Mato Grosso, thus expanding the P_{total} regional dataset prepared by Castanho et al. (2013); (2) to develop a new dataset of V_{max} from the P_{total} regional dataset and (3) to generate a second dataset of V_{max} based on a global P database (Yang et al., 2013). These V_{max} datasets will be used

in the next chapter for photosynthesis limitation in the experiment with the INLAND model.

1.2 MATERIALS AND METHODS

The methodology described in this chapter includes: (1) Description of the phosphorus regional data; (2) Estimation of P_{total} from phosphorus $P_{-Melich-1}$; (3) Development of dataset of P_{total} (PR); (4) Development of two V_{max} datasets from regional (PR) and global (PG) databases, for use by INLAND;

1.2.1 Data Description of phosphorus in soil

In Soil Science, numerous extractors that use different methodologies to extract the lability of the forms of P in the soil are commonly used. In this study, we used P data obtained via the Mehlich-1 extractor (H₂SO₄, 0.025mol L⁻¹ + HCl 0.05mol L⁻¹) 1:10 soil: solution ratio (P_{-Melich-1}), and the resin method (P_{resin}) databases to estimate total phosphorus (P_{total}).

The Mehlich-1 extractor has been performing reasonably well as an indicator of the availability of P in the soil and with the application of soluble phosphate fertilizers. The method of resin anion-exchange is an attempt to reproduce in the laboratory the P absorption process by the plants in the field. The Hedley fractionation (Hedley and Stewart, 1982) uses sequentially extracts from lower to higher desorption power and allow the separation of P forms according to their nature (organic or inorganic) and facility of desorption (Cross and Schlesinger, 1995). This method consists of a sequential extraction of reagents of increased strength.

1.2.2 Description of the phosphorus regional data

We used 54 samples of phosphorus content (P) in the soil, where 52 were obtained via the Mehlich-1 extractor, and two were direct measurements of total phosphorus content (P_{total}) in the soil. In addition to these samples collected in the field, a soil map of the Amazon was also used (Quesada et al., 2010).

The 54 samples of P content were provided by researchers group at the UNEMAT – State University of Mato Grosso, collected for different vegetation types ranging from sparse physiognomies like *Campo de Murundus*, open flooded field, *Cerrado típico* to more dense forest formations such as *Cerradão*, semideciduous forest, evergreen seasonal forest and gallery forest in the Amazon-Cerrado transition region in the state of Mato Grosso (Table 1).

The information on *Cerradão*, *Cerrado típico*, *Cerrado rupestre* and Gallery Forest are from the Bacaba Municipal Park conservation area, which is located in Nova Xavantina-MT (14° 41'S and 52° 20'W). The Bacaba Municipal Park has 489 ha of natural landscape with Cerrado vegetation type with the prevalence of *Cerrado sensu-stricto*, with medium to clayey soil texture (Marimon Junior and Haridasan, 2005). For the physiognomies open field and *Campo de Murundus*, the phosphorus samples are from the Araguaia State Park, located in the northeast of Mato Grosso State, in the municipality of Novo Santo Antonio (11° 37' 02"S and 50° 40' 10"W), inserted in the extensive sedimentary plain of Bananal in the Araguaia River basin.

The spatial distribution of these samples can be seen in Figure 1, which represents the total phosphorus in the soil with a spatial resolution of $1^{\circ} \times 1^{\circ}$ (Quesada et al., 2010).



Figure 1. Location and geographical distribution on the regional dataset P_{total} (Quesada et al., 2010) showing the areas of phosphorus sample collection of $P_{-Mehlich-1}$ in Mato Grosso state. The pink dots are locations of sampling and the thick black line is the geographical limit of the Cerrado biome.

Table 1. Phosphorus content data (P_{-Mehlich-1}) and clay percentage for each sample collected in the Amazon-Cerrado transition region.

Location		Dhysiagnamy	Phosphorus	Clay		
Latitude	Longitude	n	Physiognomy	mg kg ⁻¹	%	
-15.55	-50.10	1	Cerrado rupestre ^a	0.89	30.6	
-15.54	-50.10	2	Cerrado típico ^b	0.20	34.7	
-14.17	-51.76	3	Cerrado ralo ^c	2.28	21.1	
-14.17	-51.77	4	Cerrado ralo ^c	1.30	30.0	
-14.15	-51.76	5	Cerrado típico ^b	2.93	40.5	
-14.16	-51.77	6	Cerrado típico ^b	1.11	35.4	
-14.71	-52.35	7	Cerrado típico ^b	3.00	35.8	
-14.71	-52.35	8	Cerrado típico ^b	0.84	48.2	
-14.71	-52.35	9	Cerrado típico ^b	0.42	49.3	
-14.82	-52.17	10	Semi deciduous Forest	3.18	21.5	
-14.71	-52.35	11	Cerrado típico ^b	0.34	17.3	
-14.71	-52.35	12	Cerrado típico ^b	0.13	17.7	
-14.70	-52.35	13	$Cerrad ilde{a}o^d$	0.26	21.0	
-14.70	-52.35	14	$Cerrad ilde{a}o^d$	0.10	24.4	
-14.69	-52.35	15	$Cerrad ilde{a}o^d$	5.46	21.0	
-14.69	-52.35	16	<i>Cerradão</i> ^d	3.80	33.5	
-14.69	-52.35	17	<i>Cerradão^d</i>	1.90	40.5	
-14.69	-52.35	18	$Cerrad ilde{a}o^d$	0.80	44.0	

Location			Dhusiaanamu	Phosphorus	Clay
Latitude	Latitude Longitude n		Physiognomy	mg kg ⁻¹	%
-14.69	-52.35	-52.35 19 Cerradão ^d		0.30	45.2
-14.72	-52.36	20	Gallery Forest	0.87	15.0
-14.72	-52.36	21	Gallery Forest	6.94	10.5
-14.72	-52.36	22	Flooded Gallery Forest	1.71	11.7
-13.10	-53.39	23	Flooded Riparian Forest	26.0	43.0
-13.10	-53.39	24	Flooded Riparian Forest	18.0	49.0
-13.00	-50.25	25	Cerrado rupestre ^a	2.44	4.44
-12.38	-50.93	26	Campo de Murundus ^g	2.30	39.3
-12.36	-50.93	27	Campo de Murundus ^g	3.30	29.5
-12.56	-50.92	28	Campo de Murundus ^g	3.30	22.0
-12.04	-50.73	29	Campo de Murundus ^g	0.70	38.5
-12.57	-50.91	30	Campo de Murundus ^g	1.70	39.1
-12.62	-50.82	31	Campo de Murundus ^g	2.40	25.6
-12.43	-50.72	32	Campo de Murundus ^g	2.30	29.1
-12.22	-50.77	33	Campo de Murundus ^g	2.30	30.8
-12.37	-50.94	34	Campo de Murundus ⁸	5.20	37.5
-12.37	-50.94	35	Campo de Murundus ⁸	4.00	39.3
-12.38	-50.93	36	open flooded field	1.60	32.5
-12.38	-50.93	37	open flooded field	2.00	20.8
-12.36	-50.93	38	open flooded field	1.60	17.1
-12.04	-50.73	39	open flooded field	0.80	22.8
-12.57	-50.91	40	open flooded field	0.70	25.0
-12.62	-50.82	41	open flooded field	2.20	20.8
-12.43	-50.72	42	open flooded field	1.60	19.6
-12.22	-50.77	43	open flooded field	1.90	27.0
-12.37	-50.94	44	open flooded field	3.10	29.9
-12.37	-50.94	45	open flooded field	0.90	30.8
-12.83	-52.35	46	Seasonal Evergreen Forest	Ptotal=141	49.0
-12.81	-51.85	47	Seasonal Evergreen Forest	$\mathbf{P}_{\text{total}} = 117$	16.0
-11.18	-50.23	48	Cerrado denso ^h	2.71	3.96
-11.18	-50.23	49	$Cerrad ilde{a}o^d$	1.66	4.16
-11.17	-50.23	50	Cerrado típico ^b	1.45	3.56
-11.86	-50.72	51	open flooded field	2.00	41.6

L	ocation			Phosphorus	Clay
Latitude Longitude n		n	Physiognomy	mg kg ⁻¹	%
-11.86	-50.72	52	Campo de Murundus ^g	2.80	47.7
-9.11	-54.23	53	Cerrado rupestre ^a	4.17	4.56
-9.79	-50.43	54	Semi deciduous Forest	2.04	18.4

Table 1. (Continued.)

^{*a*}**Cerrado rupestre**: a tree-shrub vegetation that grows in areas of accentuated topography with many rock outcrops and shallow soils, where individual trees establish themselves in clefts in the rocks so that their densities will vary as a function of the specific conditions of each site (Ribeiro and Walter, 2008).

^bCerrado típico: a vegetation of trees and shrubs fairly regular and usually not exceeding 4 meters (Ribeiro and Walter, 2008).

^cCerrado ralo: a vegetation that is more open than *Cerrado típico*; the trees not exceeding 2 to 3 meters in height, covering from 5 to 20% of the soil (Ribeiro and Walter, 2008).

^dCerradão: a dense and tall woodland formation (Ribeiro and Walter, 2008).

^gCampo de Murundus: a typical landscape of Central Brazil characterized by countless rounded earth mounds (the '*murundus*'), which are covered by woody '*Cerrado*' vegetation and are found scattered over a grass-covered surface (the 'campo') (Ribeiro and Walter, 2008).

^hCerrado denso: this vegetation is more dense than *Cerrado típico*; the trees exceeding 2 to 3 meters in height, and covered with a woody cover ranging from 10 to 60% (Ribeiro and Walter, 2008).

1.2.3 Estimating Ptotal from P-Mehlich-1

The data for the phosphorus content in soil extracted by Mehlich-1 ($P_{-Mehlich-1}$) was used to estimate the total phosphorus content in the soil (P_{total}). For this, it was first necessary to establish a relationship between $P_{-Mehlich-1}$ and P_{total} .

Based on P_{-Mehlich-1} studies conducted by Neves (2000) and dos-Santos et al. (2008), a linear relationship was produced between P_{-Mehlich-1}, clay percent and P_{total} using a Hedley fractionation database for 26 sites in the Amazon (Castanho et al., 2013; Quesada et al., 2010). These 26 locations were the same used to construct the relationship between V_{max} and P_{total} by Castanho et al. (2013). The database of Quesada et al. (2010) provides data of physicochemical properties of the soil, and Hedley fractionation data with eight different fractions of phosphorus content in the soil, including: phosphorus extracted by resin (P_{resin}), inorganic bicarbonate, organic bicarbonate, inorganic sodium hydroxide, organic sodium hydroxide, hydrochloric acid,

 P_{total} and residual. The location, name of the experimental sites, value of P_{total} , P_{Resin} and clay percent used to establish the relationship between $P_{-Mehlich-1}$ and P_{total} are shown in Table A1 (Appendix Table A1).

Based on the Freire (2001) equation, the amount of phosphorus remaining in the soil, i.e., the existing amount of P in the soil (P_{rem}) was estimated for each site, based on their clay content:

$$P_{\rm rem} = 52.44 - 0.9646 \ \text{C} + 0.005 \ \text{C}^2 \qquad \qquad R^2 = 0.747 \tag{1}$$

where P_{rem} is expressed in mg L⁻¹ and C is the clay content in %. P_{rem} is the P concentration that remains in solution after shaking soil with 0.01 mol L⁻¹ CaCl₂ containing 60 mg L⁻¹ P (Alves and Lavorenti, 2006).

After obtaining of the remaining values (P_{rem}), it was necessary to estimate the phosphorus maximum adsorption capacity (CMAP) of each soil, in order to calculate how much phosphorus each soil is capable of adsorbing.

Based on significative data from several studies (Bognola, 1995; Campello et al., 1994; Fabres, 1986; Gonçalves, 1988; Ker, 1995; Moreira, 1988; Muniz, 1983; Novelino, 1999; Paula, 1993), Neves (2000) proposed Equation (2) to calculate CMAP from P_{rem}:

$$CMAP = 1816.1 - 373.72 \log P_{rem}$$
 $R^2 = 0.751$ (2)
where CMAP is expressed in mg kg⁻¹ and P_{rem} in mg L⁻¹.

Knowing the CMAP, Neves (2000) also proposed a robust model (Equation 3), which estimates how much the Mehlich-1 extractor is capable of removing of P added in each soil sample (P-Mehlich-1/PAdc). This relationship was adjusted based on 31 soil

samples data from the work of Bahia-Filho (1982), Muniz (1983), Gonçalves (1988) and Novelino (1999);

$$\frac{P_{-Mehlich-1}}{P_{Adc}} = 380.96 \text{ CMAP}^{-1.2101} \text{ R}^2 = 0.734, (p<0.001)$$
(3)

where P_{Adc} is the added dose of P in soil expressed in mg kg⁻¹.

Finally, knowing $P_{-Mehlich-1}/P_{Adc}$, $P_{resin}/P_{-Mehlich-1}$ was estimated using the Equation (4) established by Neves (2000) with r=0.899, n=26.

$$\frac{P_{\text{resin}}}{P_{-\text{Mehlich}-1}} = 0.5553 \left(\frac{P_{-\text{Mehlich}-1}}{P_{\text{Adc}}}\right)^{0.6002} \qquad R^2 = 0.808, (p<0.001) \tag{4}$$

where P_{resin} is expressed in mg kg⁻¹.

With the P_{resin} values and the ratio estimated by Equation (4), $P_{-Mehlich-1}$ values for all stations in Quesada et al. (2010) were estimated (P_{m1_est} , expressed in mg kg⁻¹). Table 2 shows the estimates obtained by Equations (1), (2), (3) and (4) for the 26 sites in the Amazon.

Obtaining $P_{m1_{est}}$ values for locations where data for clay content and P_{total} (mg kg⁻¹) were available, enabled the development of a linear regression model (Equation 5) that estimates P_{total} from $P_{m1_{est}}$ (mg kg⁻¹) and C (%) with r = 0.639.

$$P_{\text{total}} = 72.4628 + 0.639 \text{ (P}_{m1\text{_est}} \text{ C)} \qquad R^2 = 0.408, (p<0.001) \qquad (5)$$

Although the R² value is low, the regression is significant at p<0.01. The product $(P_{m1_est} C)$ was used to correct the effect of soil clay contribution on P_{-Mehlich-1} values, which tends to remove smaller amounts of P, for high values of clay. Applying the Equation (5) to the data presented in Table 1, P_{total} was estimated for all field samples collected in Mato Grosso. These results are presented in Table 3 in Section 1.3 (Results

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Observation data					Estimated of	lata	
Quesada	et al. (20	10)	D		P _{-Mehlich-1}	P _{resin}	D
Plot	Presin	Clay	Presin	СМАР	P _{Adc}	P _{-Mehlich-1}	P_m1_est
ID	mg kg ⁻¹	%	mg L ⁻¹	mg kg ⁻¹	mg kg ⁻¹	mg kg ⁻¹	mg kg ⁻¹
'RIO-12'	3.42	9.50	43.7	404.2	0.27	1.23	2.79
'ELD-12'	5.34	20.1	35.1	486.9	0.21	1.40	3.80
'SCR-01'	2.36	6.88	46.0	384.9	0.28	1.18	1.99
'TIP-05'	9.01	37.3	23.4	637.6	0.15	1.71	5.27
'JRI-01'	1.30	80.7	7.15	1081	0.08	2.51	0.52
'JAS-02'	8.79	29.1	28.6	563.1	0.18	1.56	5.64
'CAX-01'	5.17	41.8	20.9	680.6	0.14	1.79	2.88
'MBO-01'	3.14	11.5	42.0	419.1	0.26	1.26	2.49
'BNT-04'	4.82	57.7	13.4	845.2	0.11	2.10	2.30
'TAP-04'	4.50	89.3	6.18	1136	0.08	2.60	1.73
'ALP-12'	7.37	14.0	40.0	437.9	0.24	1.30	5.67
'SUC-02'	4.06	37.2	23.5	636.7	0.15	1.71	2.38
'AGP-01'	2.99	42.6	20.4	688.8	0.14	1.81	1.66
'ZAR-03'	4.81	31.1	27.3	580.7	0.17	1.60	3.01
'TAP-123'	1.65	66.1	10.5	936.4	0.10	2.26	0.73
'ZAR-04'	7.48	18.3	36.5	471.8	0.22	1.37	5.45
'JUR-01'	10.6	36.6	23.8	631.2	0.16	1.70	6.22
'RST-01'	8.29	25.4	31.2	530.8	0.19	1.49	5.55
'ALF-01'	3.51	11.4	42.1	418.7	0.26	1.26	2.79
'DOI-01'	7.18	19.1	35.9	478.5	0.22	1.39	5.18
'SIN-01'	2.58	9.8	43.5	406.4	0.27	1.23	2.10
'TAM-01'	5.85	37.8	23.2	641.9	0.15	1.72	3.41
'CUZ-03'	11.5	42.5	20.5	687.4	0.14	1.80	6.35
'CRP-01'	21.8	18.1	36.7	470.1	0.22	1.37	15.9
'HCC-21'	7.34	25.6	31.0	532.4	0.19	1.50	4.90

Table 2. Estimates made for obtaining $P_{-Mehlich-1}$ values for the stations used in this study.

1.2.4 Development and description of the phosphorus regional dataset (PR)

The 54 P_{total} values were used to extend (Table 3) the map of Quesada et al. (2010) in the Amazon-Cerrado transition. Depending on the spatial distribution of the

samples, the average for the P_{total} values (Table 4) was calculated inside each $1^{\circ} \times 1^{\circ}$ pixel and the new values added are shown in Figure 2b.



Figure 2. P_{total} regional dataset in mg kg⁻¹ (Quesada et al., 2010) (a), PR with new estimated P_{total} data (b), In both datasets the limit of the Cerrado biome is defined by the thick black line. The sites that provided additional phosphorus data are represented by pink dots.

1.2.5 Description of the global phosphorus dataset (PG)

A global dataset of total phosphorus content in the soil (P_{total}) with a spatial resolution of $0.5^{\circ} \times 0.5^{\circ}$ was also used (Figure 3), which is part of a database containing six global maps of the different forms of phosphorus found in soil that include: labile inorganic phosphorus, organic phosphorus, occluded phosphorus, secondary mineral phosphorus, phosphorus in the form of apatite and total phosphorus (P_{total}) (Yang et al., 2013). These maps were made from lithologic maps, distribution of soil development stages, fraction of the remaining source material for different stages of weathering using chronosequence studies (29 studies), and phosphorus distribution in different forms for

each soil type based on analysis of Hedley fractionation (Yang and Post, 2011) which are part of a worldwide collection of soil profile data.



Figure 3. Map of total phosphorus (mg kg⁻¹) in the soil (Yang et al., 2013).

The uncertainties and limitations of this database are related to the Hedley fractionation data used. When quantified, these uncertainties are 17% for low weathered soils, 65% for intermediate soils and 68% for highly weathered soils (Yang et al., 2013).

1.2.6 Development of V_{max} regional and global datasets

The V_{max} values in the INLAND model are defined for each plant functional type which, combined in different forms, create the distinct ecosystems represented by the model. Default V_{max} values for tropical evergreen trees are defined at 65 μ molCO₂ m⁻² s⁻¹.

Here, the V_{max} datasets were produced through a linear relationship proposed by Castanho et al. (2013), who used soil fertility (P_{total}) to limit photosynthesis in the
INLAND model by estimating the maximum capacity of carboxylation by the Rubisco enzyme (V_{max}), applying Equation (6) to the regional and global datasets of total phosphorus (PR and PG).

$$V_{\text{max}} = 0.1013 P_{\text{total}} + 30.037 \tag{6}$$

where V_{max} and P_{total} are given in μ molCO₂ m⁻² s⁻¹ and mg kg⁻¹, respectively.

1.3 RESULTS

1.3.1 Estimates of total phosphorus

Table 3 shows the total P values estimated for 54 samples of $P_{-Mehlich-1}$ in Mato Grosso state. These samples were spatialized to a grid of $1^{\circ} \times 1^{\circ}$ resolution and resulted in 12 new pixels. Due to the large size of the grid (approximately 111 km), different physiognomies were grouped into a single pixel. For each pixel, the average of P_{total} values was considered regardless of the type of vegetation.

The average values per physiognomy were higher for the Riparian Forest with $P_{total} = 11.46 \text{ mg kg}^{-1}$ (n=2) followed by *Campo de Murundus* with $P_{total} = 132.8 \text{ mg kg}^{-1}$ (n=11), Semi deciduous Forest with $P_{total} = 106.3 \text{ mg kg}^{-1}$ (n=2), *Cerradão* with $P_{total} = 103.0 \text{ mg kg}^{-1}$ (n=8), open flooded field with $P_{total} = 100.8 \text{ mg kg}^{-1}$ (n=11), *Cerrado ralo* with $P_{total} = 100.2 \text{ mg kg}^{-1}$ (n=2), Gallery Forest with $P_{total} = 99.8 \text{ mg kg}^{-1}$ (n=2), *Cerrado típico* with $P_{total} = 97.1 \text{ mg kg}^{-1}$ (n=8), Flooded Gallery Forest with $P_{total} = 85.2 \text{ mg kg}^{-1}$ (n=1), *Cerrado rupestre* with $P_{total} = 84.62 \text{ mg kg}^{-1}$ (n=3), and *Cerrado denso* with $P_{total} = 79.32 \text{ mg kg}^{-1}$ (n=1).

L	ocation			Phosphorus	Ptotal
Latitude	Longitude	n	Physiognomy	mg kg ⁻¹	mg kg ⁻¹
-15.55	-50.1	1	Cerrado rupestre ^a	0.89	89.9
-15.54	-50.1	2	Cerrado típico ^b	0.20	76.9
-14.17	-51.76	3	Cerrado ralo ^c	2.28	103
-14.17	-51.77	4	Cerrado ralo ^c	1.30	97.4
-14.15	-51.76	5	Cerrado típico ^b	2.93	148
-14.16	-51.77	6	Cerrado típico ^b	1.11	97.6
-14.71	-52.35	7	Cerrado típico ^b	3.00	141
-14.71	-52.35	8	Cerrado típico ^b	0.84	98.3

Table 3. Ptotal estimates for the 54 soil samples P-Mehlich-1 in the Mato-Grosso.

 Table 3. (Continued.)

L	Location			Phosphorus	Ptotal	
Latitude	Longitude	n	Physiognomy	mg kg ⁻¹	mg kg ⁻¹	
-14.71	-52.35	9	Cerrado típico ^b	0.42	85.7	
-14.82	-52.17	10	Semi deciduous Forest	3.18	116	
-14.71	-52.35	11	Cerrado típico ^b	0.34	76.2	
-14.71	-52.35	12	Cerrado típico ^b	0.13	73.9	
-14.7	-52.35	13	$Cerrad ilde{a}o^d$	0.26	76.0	
-14.7	-52.35	14	$Cerrad ilde{a}o^d$	0.10	74.0	
-14.69	-52.35	15	$Cerrad ilde{a}o^d$	5.46	146	
-14.69	-52.35	16	$Cerrad ilde{a}o^d$	3.80	154	
-14.69	-52.35	17	$Cerrad ilde{a}o^d$	1.90	122	
-14.69	-52.35	18	$Cerrad ilde{a}o^d$	0.80	95.0	
-14.69	-52.35	19	<i>Cerradão</i> ^d	0.30	81.1	
-14.72	-52.36	20	Gallery Forest	0.87	80.1	
-14.72	-52.36	21	Gallery Forest	6.94	119	
-14.72	-52.36	22	Flooded Gallery Forest	1.71	85.2	
-13.1	-53.39	23	Flooded Riparian Forest	26.0	787	
-13.1	-53.39	24	Flooded Riparian Forest	18.0	636	
-13	-50.25	25	Cerrado rupestre ^a	2.44	79.4	
-12.38	-50.93	26	Campo de Murundus ^g	2.30	130	
-12.36	-50.93	27	Campo de Murundus ^g	3.30	135	
-12.56	-50.92	28	Campo de Murundus ^g	3.30	119	
-12.04	-50.73	29	Campo de Murundus ^g	0.70	89.7	
-12.57	-50.91	30	Campo de Murundus ^g	1.70	115	
-12.62	-50.82	31	Campo de Murundus ^g	2.40	112	
-12.43	-50.72	32	Campo de Murundus ^g	2.30	115	
-12.22	-50.77	33	Campo de Murundus ^g	2.30	118	
-12.37	-50.94	34	Campo de Murundus ^g	5.20	197	
-12.37	-50.94	35	Campo de Murundus ^g	4.00	173	
-12.38	-50.93	36	open flooded field	1.60	106	
-12.38	-50.93	37	open flooded field	2.00	99.0	
-12.36	-50.93	38	open flooded field	1.60	89.9	
-12.04	-50.73	39	open flooded field	0.80	84.1	
-12.57	-50.91	40	open flooded field	0.70	83.6	
-12.62	-50.82	41	open flooded field	2.20	102	

Table 3. ((Continued.)

Location		-		Phosphorus	Ptotal
Latitude	Longitude	n	Physiognomy	mg kg ⁻¹	mg kg ⁻¹
-12.43	-50.72	42	open flooded field	1.60	92.5
-12.22	-50.77	43	open flooded field	1.90	105
-12.37	-50.94	44	open flooded field	3.10	132
-12.37	-50.94	45	open flooded field	0.90	90.2
-12.83	-52.35	46	Seasonal Evergreen Forest		142
-12.81	-51.85	47	Seasonal Evergreen Forest		117
-11.18	-50.23	48	Cerrado denso ^h	2.71	79.3
-11.18	-50.23	49	$Cerrad ilde{a}o^d$	1.66	76.9
-11.17	-50.23	50	Cerrado típico ^b	1.45	75.8
-11.86	-50.72	52	Campo de Murundus ^g	2.80	158
-11,86	-50,72	51	open flooded field	2.00	126
-9.11	-54.23	53	Cerrado rupestre ^a	4.17	84.6
-9,79	-50,43	54	Semi deciduous Forest	2.04	96.4

^{*a*}**Cerrado rupestre**: a tree-shrub vegetation that grows in areas of accentuated topography with many rock outcrops and shallow soils, where individual trees establish themselves in clefts in the rocks so that their densities will vary as a function of the specific conditions of each site (Ribeiro and Walter, 2008).

^{*b*}**Cerrado típico**: a vegetation of trees and shrubs fairly regular and usually not exceeding 4 meters (Ribeiro and Walter, 2008).

^cCerrado ralo: a vegetation is more open than *Cerrado típico*; the trees not exceeding 2 to 3 meters in height, covering from 5 to 20% of the soil (Ribeiro and Walter, 2008).

^dCerradão: a dense and tall woodland formation (Ribeiro and Walter, 2008).

^gCampo de Murundus: are a typical landscape of Central Brazil characterized by countless rounded earth mounds (the '*murundus*'), which are covered by woody '*Cerrado*' vegetation and are found scattered over a grass-covered surface (the 'campo') (Ribeiro and Walter, 2008).

^hCerrado denso: this vegetation is more dense than *Cerrado típico*; the trees exceeding 2 to 3 meters in height, and covered with a woody cover ranging from 10 to 60% (Ribeiro and Walter, 2008).

1.3.2 Main differences between phosphorus datasets used in this study

Figure 4 shows the difference of the soil phosphorus content between the global

P_{total} map (PG) and the regional map (PR). It is possible to observe that the global data

underestimates the Ptotal values in some Amazon-Cerrado transitional areas, mainly in

western Amazonia. PG overestimations are observed in northern of Amazonia and in

most of the Cerrado biome area.



Figure 4. Difference between the global P_{total} maps and regional P_{total} map (PG-PR) in mg kg⁻¹. The Cerrado biome is delimited by the thick black line.

The differences between the absolute values of total phosphorus at a spatial resolution of $1^{\circ} \times 1^{\circ}$ varied in the range of $\pm 180 \text{ mg kg}^{-1}$, with an average of 24.19 mg kg⁻¹.

1.3.3 V_{max} maps used in this study

Equation 6 was applied to the regional map of P_{total} (Section 1.2.6) to obtain the regional V_{max} map (PR), and to the global P_{total} map (Section 1.2.5) to generate the global V_{max} map (PG). Figure 5 shows V_{max} calculated from the P datasets. In Figure 5a, all pixels received the same value for V_{max} (65 µmolCO₂ m⁻² s⁻¹), corresponding to the default settings of INLAND. In Figures 5b and 5c it is possible to observe that the V_{max} data followed the spatial gradient of the total P content in soil, with values varying between 36 and 76 µmolCO₂ m⁻² s⁻¹.



Figure 5. Default V_{max} value of INLAND – PC (a), Regional V_{max} dataset – PR (b), Global V_{max} dataset – PG (c), all in μ molCO₂ m⁻² s⁻¹.

The estimated values of P_{total} in the Amazon-Cerrado border were used to expand the regional P map used by Castanho et al. (2013) and thus generate the V_{max} map shown in Figure 5b. These new values are described in Table 4 and range from 37.9 to 102.1 μ molCO₂ m⁻² s⁻¹.

Samples		Average P _{total}	V _{max}
n=54	Pixel	mg kg ⁻¹	µmolCO ₂ m ⁻² s ⁻¹
2	1	83.38	38.48
4	2	111.6	41.34
16	3	101.5	40.32
2	4	711.5	102.1
1	5	79.39	38.08
20	6	114.3	41.62
1	7	141.5	44.38
1	8	117.0	41.89
3	9	77.32	37.87
2	10	141.7	44.39
-	11	84.61	38.61
1	12	96.40	39.80

Table 4. V_{max} values and P_{total} values for samples in the Amazon-Cerrado transition region.

1.4 CONCLUSIONS

The recent availability of phosphorus data in the Amazon made possible the development of a model able to estimate P_{total} throught $P_{-Mehlich-1}$, thus extending the regional soil map in the Amazon-Cerrado transition. This methodology allows new estimates of P_{total} to be obtained from $P_{-Mehlich-1}$ data collected in the field, using a robust model with 99% confidence in the estimates.

Although a model was developed to estimate P_{total} in the soil, it is recommended that future works interested in evaluating the P content in the plants should not collect only the labile P, but use the Hedley fractionation method, which informs all forms of P for the given soil sample.

The results produced by the estimates of P_{total} were incorporated into the total phosphorus regional dataset, generating a new map of regional V_{max} . The V_{max} maps produced in this work are potentially useful tools to DGVMs in general, and are used in the numeric experiment in the next chapter.

CHAPTER 2 – Effects of climate, fire and phosphorus in the Amazon-Cerrado border simulated with INLAND model – Numeric experiment development

2.1 INTRODUCTION

The transition between the Amazon and Cerrado in Brazil has the largest area of contact between forest and savanna in the tropical regions and these biomes differ fundamentally in their structural characteristics and species composition (Torello-Raventos et al., 2013). In this transition, climate seasonality – especially precipitation – and fire disturbances have an important role in the balance of the vegetation due to influences of the ecological and biogeochemical processes of vegetation directly affecting the carbon fluxes (for example, the Gross Primary Production - GPP, Net Primary Production – NPP and respiration) that, over time, may lead to changes in composition and structure of vegetation (Alves et al., 1997). Tree species associated with forest or savanna vegetation differ in numerous physiological characteristics, such as fire survivorship (Hoffmann et al., 2009; Ratnam et al., 2011), as well as in their wood and foliar characteristics (Gotsch et al., 2010), sometimes related with soil fertility (Ferreira-Ribeiro and Tabarelli, 2002; Miranda et al., 2003; Quesada et al., 2012). Generally, we have an incomplete knowledge on how the trees of these vegetation types differ in photosynthesis characteristics, especially in terms of response to nutrient availability.

In the transition Amazonia-Cerrado there are no studies assessing the influence of factors such as nutritional limitation by P in the photosynthesis rates of woody trees. However, studies that quantify the P levels in soils along the Amazon rainforest in its various forms contribute to information about the P content in the soil (Quesada et al., 2009; Quesada et al., 2010; Quesada et al., 2012; Yang et al., 2013). In Cerrado regions, the quantification of phosphorus is poor, since it was believed that, in contrast to tropical forests, photosynthetic capacity in the savannas would be more likely to be limited by N than by P (Lloyd et al., 2009).

However, works from West Africa showed that, depending on the relative concentrations of P and N in the leaves, both Rubisco activity and electron transport activity of African savanna and forest trees can potentially be limited by either N or P (Domingues et al., 2010), so there is no consensus about how the vegetation is limited by these nutrients and how this limitation associated with factors such as climate and fire could act on the areas of transition in a scenario of future climate change. Numerous efforts through dynamic vegetation models (DGVMs) have been conducted on Amazonia forests with the aim of understanding dynamics through the analysis of long-term field observations of tree growth and mortality patterns as they relate to climatic and edaphic variations across the basin (Castanho et al., 2013; Phillips et al., 2004; Quesada et al., 2012).

The INLAND model is a DGVM that has been improved by the Brazilian scientific community to better represent intrinsic characteristics of tropical biomes like fire and nutritional limitation. The objective of this chapter is to simulate the current position of the Amazon-Cerrado border, considering different combinations among nutritional limitation, occurrence of fire, interannual climate variability and mean climate to investigate the influence of these factors on the dynamics of the Amazonia-Cerrado transition. The specific objectives are 1) generate a leaf area index (LAI) map for the Amazonia-Cerrado transition for validation of the simulations, 2) compare LAI values and biomass simulated by the model with the values of leaf area index and biomass from MODIS (MOD15A2 product) in five longitudinal transects in the

2.2 MATERIALS AND METHODS

2.2.1 Description of the INLAND Surface Model

The INLAND model is the main and the first land-surface component of the Brazilian Earth System Model (BESM). INLAND is based on the IBIS model (Foley et al., 1996) which considers changes in the composition and structure of vegetation in response to the environment and incorporates important aspects of biosphereatmosphere interactions in South America. The model simulates the exchanges of energy, water, carbon and momentum between soil-vegetation-atmosphere. These processes are organized in a hierarchical framework and operate at different time steps, ranging from 60 minutes to 1 year, coupling ecological, biophysical and physiological processes (Kucharik et al., 2000). The vegetation is represented by two layers, upper and lower canopy, and 12 plant functional types (PFTs): tropical evergreen trees, tropical deciduous trees, temperate evergreen trees (broadleaf), temperate evergreen trees (conifers), temperate deciduous trees (broadleaf), boreal coniferous evergreen trees, boreal deciduous trees (broadleaf), boreal deciduous trees (coniferous), perennial shrubs, deciduous shrubs, herbaceous C3 and grasses C4. The combination of the 12 PFTs creates the distinct ecosystems represented by the model: tropical evergreen forest, tropical deciduous forest, temperate evergreen broadleaf forest, temperate evergreen conifer forest, temperate deciduous forest, boreal evergreen forest, boreal deciduous forest, mixed forest, savanna, grassland, dense shrubland, open shrubland, tundra, desert and polar desert or ice.

The leaf photosynthesis and respiration are calculated used Farquhar equations (Farquhar et al., 1980; Sharkey, 1985), where the gross photosynthesis rate per unit leaf area (Ag) is a function of absorbed light, leaf temperature, internal carbon dioxide

concentration, and the Rubisco enzyme capacity for photosynthesis (V_{max}). The maintenance respiration is a function only of V_{max} . The yearly dynamic vegetation module computes the following for each PFT: gross and net primary productivity (GPP and NPP), changes in biomass pools, simple mortality disturbance processes and resultant Leaf Area Index (LAI), thus allowing vegetation type and cover to change through time. GPP and NPP are calculated at the end of each year as:

$$GPP = \int Ag_{dti} \tag{7}$$

$$NPP = (1 - \eta) \int (Ag, i - R_{leaf,i} - R_{wood,i} - R_{root,i}) dt$$
(8)

where η (0.33) is the fraction of carbon lost in the construction of net plant material because of growth respiration (Amthor, 1984). The partitioning of the NPP for each plant functional type resolves carbon in three biomass pools: leaves, stems and fine roots. The leaf area index (LAI) of each PFT is obtained by simply dividing leaf carbon by specific leaf area, which in INLAND is considered fixed (one value) for each PFT.

The soil physical properties in INLAND used a multilayer formulation of soil (eight soil layers) to simulate the diurnal and seasonal variations of heat and moisture. Each layer is described in terms of soil temperature, volumetric water content and ice content (Foley et al., 1996; Thompson and Pollard, 1995). Furthermore, all of these processes are influenced by these soil texture and amount of organic matter within the soil profile.

Considering these aspects vegetation dynamics and soil physical properties the model can simulate plant competition for light and water between trees, shrubs and herbaceous through shading and differences in water uptake (Foley et al., 1996).

The soil chemical properties are represented by the carbon cycle (C), nitrogen (N) and phosphorus (P), the latter being recently implemented by Castanho et al. (2013). The carbon cycle is simulated through vegetation, litter and soil organic matter, where the biogeochemical module is similar to the CENTURY model (Parton et al., 1993; Verberne et al., 1990). The amount of C existing in the first meter of soil is divided into different compartments characterized according to their time residence, which can vary in an interval of a few hours for microbial biomass and organic material to several years for lignin. For the nitrogen, the model considers only the soil N transformations and carbon decomposition, not influencing the productivity of vegetation, ie., there is a fixed C:N ratio. Phosphorus is used only to limit the net primary productivity. The total phosphorus available in the soil is used to estimate the maximum capacity of carboxylation by the enzyme Rubisco (V_{max}) through a linear relationship, Equation 6.



Figure 6. INLAND model structure.

INLAND also contains two fire modules. The first module is a simple fixedvalue disturbance, which does not depend on the environmental conditions. The second module is based on the fire module of the Canadian model of fire CTEM (Arora and Boer, 2005). In this module, all three aspects of the fire triangle – the availability of fuel to burn, the flammability of vegetation depending on environmental conditions, and the presence of an ignition source – are taken into account. Here, CTEM uses an arbitrary anthropogenic fire probability which is summed to the natural ignition probability.

2.2.2 Model Configuration and Experiment Design

The model was forced with a prescribed climate based on the Climate Research Unit (CRU) databases of the University of East Anglia (New et al., 1999). Two boundary conditions were used; the first is referred to as the monthly climatological average (CA), commonly used by the scientific community to represent the average climate of a 30 year period of the last century (1961-1990). The second boundary condition is the historical dataset, used to represent the actual climate variability for the period 1948-2008 (CV), a continuous record of 61 years. The dataset has a 1-degree spatial resolution and a monthly time resolution.

Soil texture data is based on the IGBP-DIS global soil (Global Soil Data Task 2000) (Hansen and Reed, 2000), and the Quesada et al. (2010) dataset. All of the simulations started with the potential vegetation defined by the Center for Sustainability and the Global Environment (SAGE)(Ramankutty and Foley, 1999), with soil carbon and litter pools set to zero.

In the experiment design, the control simulations (PC) to both climate scenarios (CA and CV) considered lack of nutritional limitation per phosphorus, CO₂ fixed at

380 ppm and no fire effects (fire module off), for a period of 428 years (equivalent to the period 1581-2008). In the control run (PC) the V_{max} data used to calculate productivity were the default global value used by the dynamic vegetation module of the INLAND (65 µmolCO₂ m⁻² s⁻¹) for the plant functional types: tropical evergreen trees and tropical deciduous trees. Simulations with different combinations between the climate scenarios (CA and CV), different nutritional limitation scenarios provided by V_{max} maps (PR and PG), fire effects (F) on or off, and the effects of CO₂ fixed (380 ppm) or variable CO₂ (270 ppm to 385 ppm) were also performed. A summary of the simulations are presented in Tables 5 and 6.

Table 5. Simulations with different combinations of the factors evaluated by INLAND model considering the climate scenario CA, climatological average (1961-1990).

CO	Fina (F)		Vmax				
	File (F)	PC	PR	PG			
Fixed	Off	CA+PC	CA+PR	CA+PG			
Fixed	On	CA+PC+F	CA+PR+F	CA+PG+F			
Variable	Off	CA+PC+CO ₂	CA+PR+CO ₂	CA+PG+CO ₂			
Variable	On	CA+PC+F+CO ₂	CA+PR+F+CO ₂	CA+PG+F+CO ₂			

Table 6. Simulations with different combinations of the factors evaluated by INLAND model considering the climate scenario CV, 1948-2008 monthly data.

CO.	Eine (E)		V _{max}		
002	rire (r)	PC	PR	PG	
Fixed	Off	CV+PC	CV+PR	CV+PG	
Fixed	On	CV+PC+F	CV+PR+F	CV+PG+F	
Variable	Off	$CV+PC+CO_2$	CV+PR+CO ₂	$CV+PG+CO_2$	
Variable	On	CV+PC+F+CO ₂	CV+PR+F+CO2	$CV+PG+F+CO_2$	

These combinations of the different nutritional limitation scenarios (V_{max}), climate (CA or CV, with or without variations of CO₂) and fire (F), allow the evaluation of individual and combined effects of climate, soil chemistry, and the incidence of fire

in the NPP, biomass and LAI, and their consequent effect on the position of the Amazon-Cerrado border.

2.2.3 Analysis of the Model Sensitivity

Analyses were performed for the individual effects of climate, phosphorus, fire and CO₂ on the explanatory variables: Net Primary Production (NPP), biomass, and leaf area index of the upper and lower canopy (LAI_{upper}, LAI_{lower}).

Analysis of the isolated effect of climate variability was performed using a combination of CV+PC and CA+PC, so that the subtraction between the simulations presents the effect of climate variability isolated: $(CV+PC) - (CA+PC) = (CV-CA)_{|PC}$; in other words, this is the effect of climate variability for default V_{max} (PC).

The same logic was applied to isolate factors such as fire and phosphorus in different climate scenarios, e.g. the isolated effect of fire with an average climate scenario without influence of nutritional limitation is calculated by the difference between CA+PC+F and CA+PC, so that $(CA+PC+F) - (CA+PC) = F_{|CA PC}$. The isolated effect of fire with a climate variability scenario without influence of nutritional limitation is calculated by the difference between CV+PC+F and CV+PC, so that $(CV+PC+F) - (CV+PC) = F_{|CV PC}$.

The different combinations of climate scenarios with or without the fire effect and different phosphorus limitations are described in the Table 7. After the analysis of the individual effects, the combined effect was analyzed following the same reasoning, as shown in column E of Table 7.

	Combined Effects									
	Α	В	С	D	Ε					
	Climate	Phosphorus (P)	Fire (F)	CO ₂	CO ₂ +F					
1	(CV+PC)-(CA+PC)	(CA+PR)-(CA+PC)	(CA+PC+F)-(CA+PC)	(CA+CO ₂ +PC)-(CA+PC)	(CA+PC+F+CO ₂)-(CA+PC)					
2	(CV+PR)-(CA+PR)	(CV+PR)-(CV+PC)	(CV+PC+F)-(CV+PC)	(CV+CO ₂ +PC)-(CV+PC)	(CV+PC+F+CO ₂)-(CV+PC)					
3	(CV+PG)-(CA+PG)	(CA+PG)-(CA+PC)	(CA+PR+F)-(CA+PR)	(CA+CO ₂ +PR)-(CA+PR)	(CA+PR+F+CO ₂)-(CA+PR)					
4		(CV+PG)-(CV+PC)	(CV+PR+F)-(CV+PR)	(CV+CO ₂ +PR)-(CV+PR)	(CV+PR+F+CO ₂)-(CV+PR)					
5			(CA+PG+F)-(CA+PG)	(CA+CO ₂ +PG)-(CA+PG)	(CA+PG+F+CO ₂)-(CA+PG)					
6			(CV+PG+F)-(CV+PG)	(CV+CO ₂ +PG)-(CV+PG)	(CV+PG+F+CO ₂)-(CV+PG)					
			Effects							
	Α	В	С	D	Ε					
1	C (V-A) _{PC}	P (R-C) _{CA}	$F_{ CAPC }$	CO _{2 CA PC}	$CO_2 + F_{ CA PC}$					
2	C (V-A) _{PR}	P (R-C) _{CV}	$F_{ CV PC}$	CO _{2 CV PC}	$CO_2 + F_{ CV PC}$					
3	C (V-A) _{PG}	P (G-C) _{CA}	$F_{ CAPR }$	CO ₂ CA PR	$CO_2 + F_{ CA PR}$					
4		$P(G-C)_{ CV }$	$F_{ CV PR}$	CO _{2 CV PR}	$CO_2 + F_{ CVPR }$					
5			$F_{ CAPG}$	$CO_{2 CA PG}$	$CO_2 + F_{ CA PG}$					
6			$F_{ CV PG}$	CO _{2 CV PG}	$CO_2 + F_{ CV PG}$					

Table 7. Individual and combined effects for each simulation.

2.2.4 Determination of the best model configuration

The model configuration performace of each to determine the Amazonia-Cerrado border was evaluated using two different metrics: The first metric is the correlation between total dry season LAI simulated by the model and the MODIS dry season LAI product in five longitudinal transects along the Amazon-Cerrado border, (described in Section 2.2.5). The second metric is correlation between simulated biomass and biomass data provided by Nogueira et al. (2014). The value of biomass was estimated by matching vegetation classes mapped at a scale of 1:250 000 and 29 biomass means from 41 published studies for vegetation types classified as forest (n=2317 plots) and as either non-forest or contact zones (1830 plots). For more details, see Nogueira et al. (2014).

2.2.5 Development of the MODIS LAI Map

The MODIS product used in this work is the MOD15A2, which is responsible for providing leaf area index data every 8 days, totalizing 46 images per year. When drawing up the MODIS LAI map, nine images per year were used, corresponding to the months of July, August and September, for the period of 2000-2008, amounting to 81 leaf area index images. These images were filtered by annual coverage maps and land use also provided by MODIS through the MOD12Q1 product in order to delete information from areas of non-natural vegetation. The filter was made considering a spatial resolution of 1 km.

The map of soil cover used uses a global land cover classification by The International Global Biosphere Programme (IGBP) (Belward et al., 1999; Friedl et al., 1999; Scepan, 1999) with 16 cover classes. The LAI information were filtered out for the classes: cropland cover (12), urban and built-up (13), cropland/natural vegetation mosaics (14) and sparsely vegetated barrens (15). Then bi-linear interpolation to onedegree resolution was performed and the average of all images resulted on the average dry season leaf area index MODIS (LAI_{dry}), shown in Figure 7.



Figure 7. Map of the average dry season (July, August and September) leaf area index of MODIS product MOD15A2 from 2000 to 2008 and the five transects used in this study.

2.2.6 Statistical Analyses

The simulated variables were averaged for the last 10 years of simulation (1999-2008) and compared to LAI_{dry} and biomass from Nogueira et al. (2014) within a grid cell. For statistical comparison we used the correlation coefficient and the Student t-test to verify a significant difference at p<0.05 of each of these variables for all of the simulated area.

2.3 RESULTS AND DISCUSSION

2.3.1 Comparison of simulations (Isolated Effects)

In this section we present the results of the isolated effects of climate, phosphorus, fire and CO_2 in the NPP, biomass and LAI_{lower} and LAI_{upper} canopies.

2.3.1.1 Climate effects

Figure 8 shows the influence of the average climate (CA) and interannual variability (CV) on NPP, biomass, and LAI_{upper} and LAI_{lower}.

For both climate conditions used (CA and CV), the INLAND represents well the Amazon-Cerrado transition (Figure 8a and 8b) with higher NPP values in the forest region and lower values in Cerrado.

NPP estimates from INLAND for the Amazon forest were compared against field data from LBA project and RAINFOR network sites (from KM67, ZF-2, UFAC, AGP, CAX, TAM and ZAR) described in Table 8.

INLAND simulated NPP values between 1.26 kg-C m⁻² yr⁻¹ and 0.94 kg-C m⁻² yr⁻¹ for Brazilian sites (KM67, ZF-2, UFAC, and CAX) and NPP values up to 1.42 kg-C m⁻² yr⁻¹ for the other sites. The differences between observed and simulated data for NPP in KM67 and ZF-2 were 9% in KM67 and 1% in ZF-2 when considering CA. When climate variability was considered the differences ranged between -4% and -24%. In other sites in the west Amazon basin such as AGP and ZAR (Colombia) NPP was overestimated in CA+PC (32% and 10%) and underestimated when CV was considered (-22% and -1%). UFAC and TAM, localized in the southwest Amazon basin, showed underestimates by INLAND when using CA (-5% and -7%) and overestimates when using CV (13.9% and 20.8%).

Site	Period	Observed	Average observed	NPP IN	ILAND	Error %	Error %
		NPP NPP CA+PC C		CV+PC	CA+PC	CV+PC	
KM67	2001	1.23	1 1 4	1.24	0.04	0	-24
	2004	1.06	1.14	1.24	0.94	9	-11
ZF-2	2001	1.06	1 21	1 22	1 1 1	1	4
	2002	1.36	1.21	1.22	1.11	1	-18
UFAC	2001	1.34	1 22	1.26	1 16	5	-14
	2002	1.30	1.32	1.20	1.10	-5	-11
CAX	2004-2006	1.40	1.40	1.22	1.10	-13	-21
TAM	2005	1.53	1.53	1.42	1.27	-7	-17
ZAR	2004-2006	0.93	0.93	1.23	1.13	32	22
AGP	2004-2006	1.15	1.15	1.26	1.14	10	-1

Table 8. NPP estimates from the INLAND and the relative errors for the KM67, ZF-2, UFAC, CAX, TAM, ZAR and AGP sites.



Figure 8. Climate variability effect on NPP, biomass, LAI_{upper} and LAI_{lower} . The hatched areas indicate that the variables are statistically different compared to the control simulation at the level of 95% according to the t- test and the thick black line is the geographical limit of the Cerrado biome.

In the simulation under variable climate, NPP decreased between 0.2 and 0.4 kg-C m⁻² yr⁻¹ in biomes Amazonia and Cerrado and it was significantly different for the most part of the simulated area (Figure 8c). The values of NPP in Amazonia range

from 1.2 to 0.8 kg-C m⁻² yr⁻¹ with a northwest-southeast gradient in Amazonia, and in Cerrado range between 0.2 and 0.8 kg-C m⁻² yr⁻¹ (Figure 8). In percentage terms, NPP decreases 11% in the Amazon and 15% in the Cerrado, when the climate variability was considered.

The decrease in NPP is a consequence of the higher frequency of extreme precipitation, temperature and radiation associated with interanual climate variability, causing differences in the photosynthesis and annual carbon balance. In the Cerrado, the differences between CA and CV were higher because in this biome the seasonal variability of climate affects more intensely the vegetation, exposing the trees to soil water stress, decreasing the NPP more than in the Amazon region, where the seasonality of climate is less pronounced.

The small difference between total NPP values in CA and CV is probably related to the fact that there is offsetting between the LAI_{upper} and LAI_{lower}, which causes the total carbon fluxes to vary little when only the climate varies (Figure 8). This result confirms the findings of Botta and Foley (2002), who found low sensitivity of total LAI and NPP simulated by the IBIS model to interannual climate variability. LAI is an important variable in the INLAND model because it is the main variable used to determine the dominant vegetation type that is assigned to a pixel. The simulated LAI follows the NPP gradient, with values ranging approximately from 5 to 11 m² m⁻² for LAI_{upper} in the Amazon and 0 to 6 m² m⁻² in the Cerrado (Figure 8g and 8h). For LAI_{lower} these values range between 0 and 6 m² m⁻² in Cerrado and close to zero in Amazonia (Figure 8j and 8k).

The high values of LAI_{upper} and the low values of LAI_{lower} clearly demonstrates the competition between trees and grasses in the model. In the forest-cerrado transition it can be observed predominance of trees instead of grasses, except for the western region of the state of Mato Grosso, where higher LAI_{lower} values can be observed (Figures 8j and 8k). In the central region of the Cerrado, a small area (along 45°W) with relatively high LAI_{upper} values (Figures 8g and 8h) and low values LAI_{lower} (Figures 8j and 8k) are found, featuring a forest in both climate scenarios.

The LAI simulated values are qualitatively representative, but they are probably overestimated. In the Amazon, LAI values generally vary from 4 to 8 m² m⁻² (Carswell et al., 2002; McWilliam et al., 1993) and they are, in general, larger than the LAI observed in Cerrado. According to Miranda et al. (1997) and Hoffman et al. (2005), observed LAI in Cerrado range from below 1 to 2.5 m² m⁻². In the Amazon region, the CV LAI_{upper} decreased 11.9%, and 12.5% in the Cerrado with respect to the CA simulation, while for the LAI_{lower}, it increased 37% in the Amazon and 548% in Cerrado. The increase of 548% of LAI_{lower} results from areas dominated by very low LAI_{lower} values in CA (ie. areas between 10° S and -8° S in Cerrado domain, Figure 8k) and increased in CV to 6 - 7 m² m⁻².

The biomass values in CV were between 7 and 15 kg-C m⁻² in the Amazon, and between 1 and 7 kg-C m⁻² in the Cerrado region (Figure 8). These values follow the same spatial gradient of the NPP and LAI, with higher values in the northwest and lower values in the southeast, with significant differences between the two simulation in the entire region. In the transition, the difference between CA and CV of the annual biomass resulted in a decline between 2 and 3 kg-C m⁻². The maximum decrease was in the Northern Amazonia (4 kg-C m⁻²). In the Cerrado, some points were also influenced by this variability, reducing the upper canopy biomass by approximately 5 kg-C m⁻². In the whole Amazon, when CV was used a reduction os arboreal biomass of 10.7% was found, while in the Cerrado, there was an increase of 5%.

2.3.1.2 Phosphorus effect

The analysis of the phosphorus effect under variable climate indicates that the carbon assimilation was strongly limited by the phosphorus availability in the Amazon and the reduction in the NPP was significant for most of the Amazon area for both PR and PG datasets with respect to the PC control simulation (Figures 9 and 10). The decrease in NPP was approximately 0.2 and 0.4 kg-C m⁻² yr⁻¹ for PR and PG respectively, when compared to PC, except a few pixels in northeastern Amazonia where NPP decreased by more than 1 kg-C m⁻² yr⁻¹ for PG simulations.

In Cerrado, when PR was considered, the decreases in the NPP range from 0.1 to 0.2 kg-C m⁻² yr⁻¹ and were statistically different only in the new pixels added to the phosphorus regional dataset, as expected (Figure 9a).

In contrast, we found a small decrease in NPP for most of the Cerrado area when we considered PG. Although the decrease is not statistically significant, except for one pixel close to 52° W, it influences the decline on biomass in Cerrado (Figure 10b).

In INLAND, the NPP is directly affected by V_{max} , which is strictly dependent on soil phosphorus, and this relationship helps understand the results found. In the Amazon, the difference between the average V_{max} (PG) and average V_{max} (PR) was 0.06%. In the Cerrado, on the other hand, the magnitude V_{max} (PG) is 75% of V_{max} (PR), but the two simulations did not differ in mean stocks of biomass and NPP. This similarity between PR and PG may be due to the fact that representation of vegetation in the INLAND model in this biome is dominated by grass, which is not influenced by P in the model.

The spatial variability of LAI_{upper} (Figures 9c and Figure 10c) and biomass (Figures 9b and Figure 10b) for PR and PG reflect the spatial patterns of NPP along the Amazon-Cerrado transition.

The biomass decreased in the northwest of Amazonia and in the south (between Amazonas, Pará and northern of Mato Grosso states) from 2 to 5 kg-C m⁻² (Figure 9b) when PR was considered. The biomass gradient shows that the regional nutritional limitation is strongly influenced the boundary between Amazonia-Cerrado, following the trend of a more weathered area poor in soil P in the southeastern Amazon (Quesada et al., 2009).

On the other hand, when PG was used, a higher decrease in biomass was shown in Central Amazonia, in the northeast of Pará and in the northeast of Mato Grosso (Figure 10b), which agrees with the study of Mendes et al. (2013), which demonstrated that there is a close relationship between carbon assimilation and leaf phosphorus, in this region. Although the datasets have presented a different spatial gradient, both revealed that phosphorus limitation is higher in Amazonia than in the Cerrado.

The LAI_{upper} follows a similar pattern of the biomass (Figures 9c and 10c), since the LAI values were determined from the produced leaf biomass and the leaf specific area for each PFTs in the model. The LAI_{lower} from the simulated values of PR were not different from the simulation PC control in Cerrado, whereas the nutritional limitation map in PR did not include the Cerrado domain (Figure 9d).



Figure 9. Regional Phosphorus effect (PR) on NPP (a); biomass (b); LAI_{upper} (c); and LAI_{lower} (d). The hatched areas indicate that the variables are significantly different compared to the control simulation at the level of 95% according to the t- test and the thick black line is the geographical limit of the Cerrado biome.

The small differences found in LAI_{lower} between PG and PC (Figure 10d) could be due to the PG limited the photosynthetic rates of the arboreal component and it provided a small advantage of the water availability, favoring the increase of biomass and LAI_{lower} .



Figure 10. Global Phosphorus effect (PG) on NPP (a); biomass (b); LAI_{upper} (c); and LAI_{lower} (d). The hatched areas indicate that the variables are significantly different compared to the control simulation at the level of 95% according to the t- test and the thick black line is the geographical limit of the Cerrado biome.

2.3.1.3 Fire Effect

Overall, fire has an effect decreasing Cerrado biomass, decreasing upper canopy LAI and allowing the increase of the lower canopy LAI (Figure 11). Small parts of the dry forest also suffered the same effect, mainly in Rondonia, Bolivia and southern Peru.

The isolated effect of fire has similar behavior in CV simulations ($F_{|CV PC}$, $F_{|CV PR}$ and $F_{|CV PG}$) and strongly affected the biomass, LAI_{upper} and LAI_{upper} at the transition area (Figure 11). All these simulations resulted in decreases in biomass between 1 and 5 kg-C m⁻² along the transition, decreases in LAI_{upper} between 2 and 4 m² m⁻², and

increases in LAI_{lower} between 1 and over 5 m² m⁻². In Mato Grosso state, where transition is longest the increase of LAI_{lower} was above 4 m² m⁻² which indicates high fire activity.

In the Amazon, the small decrease of arboreal biomass in simulations reinforce the idea that the Amazon rainforest is naturally impervious to fire, contrary to the tropical savannas, which are naturally influenced by this disturbance (Lehmann et al., 2011; Murphy and Bowman, 2012). This influence of fire in tropical savannas has a great ecological importance and has been, for example, associated with the expansion of savannas in drier climate conditions as a major contributing factor (Couto-Santos et al., 2014; Silvério et al., 2013). Our results show that the fire effect in the Cerrado domain along the transition implies significant increases in LAI_{lower}, that represents shrubs and herbaceous vegetation (Figure 11g, Figure 11h, Figure 11i) and decreases significantly in LAI_{upper} (Figure 11d, Figure 11e, Figure 11f), that represents trees, thus influencing the dynamics of upper and lower canopies. Under conditions of a future climate change scenario, this competition between trees and grasses can lead to a change in vegetation structure, favoring the expansion of grasses (Malhi et al., 2009).

In the Amazon domain along the transition it is evident that, although there was a significant decrease of the biomass (Figure 11a, Figure 11b, Figure 11c) in response to fire, the decrease in leaf area index was not of the same proportions (Figure 11d, Figure 11e, Figure 11f). This may be due to the much smaller time needed to build a leaf (< 1 year), when compared to the typical interval between disturbances in the forest region (several years).

The differences in the fire effects between the Amazon and Cerrado domains may be due the fact that the Amazon has denser forests and more humidity in comparison to the Cerrado domain, and the latter is dominated by the presence of a continuous grassy layer that regains its flammability very quickly after burning. The Amazon is less flammable, generally burns less frequently and with less intensity, which allow this biome to maintain a dense canopy with distinct ecosystem properties (Eldridge et al., 2011). In the transition, the fire effects on LAI_{upper} and LAI_{lower} were inversely related. The decrease in the LAI_{upper} led to an increase in radiation available to the lower canopy, which led to the spread of a shrub and herbaceous component. Therefore, there was a significant increase in LAI_{lower}, ranging up to 5 m² m⁻² in many places in the Cerrado.



Figure 11. Fire effect on biomass, LAI_{upper} , and LAI_{lower} for all soil phosphorus. The hatched areas indicate that the variables are significantly different compared to the control simulation at the level of 95% according to the t- test and the thick black line is the geographical limit of the Cerrado biome.

2.3.2 Longitudinal gradient of biomass and leaf area index in the Amazonia-Cerrado transition – Field observations

The five longitudinal transects used in the transition's analysis showed higher values in the west and lower values in the east for biomass (Nogueira et al., 2014) and dry season MODIS leaf area index (LAI_{dry}) (Figure 12). This gradient agrees with the trends observed by Malhi et al. (2006), who found higher biomass in the central Amazon compared to lower values in the southeast Amazon region. Observed data fields in the transition region are limited, and the existing data are estimates based on small samples, which reduces in part its reliability. On the other hand, remote sensing data such as MODIS are an alternative to large-scale studies because they provide good estimates of LAI, contributing to information on the dynamics and structure of vegetation.

In general, the MODIS LAI_{dry} showed good correlation with the biomass data of Nogueira et al. (2014) for most transects analyzed, with correlation (R) values equal to 0.65 (transect 1 - T1), 0.74 (transect 2 - T2), 0.95 (transect 3 - T3), 0.89 (transect 4 - T4), and 0.54 (transect 5 - T5). The correlation between LAI_{dry} and biomass was lower in the T1 and T5 than in transects located in the central transition. In T1 this may be due to higher precipitation levels in the region, allowing the maintenance of a forest with about the same LAI along the transect, although gradually changing to a less woody structure from west to east (Figure 12). The lowest correlation between the observed data was 0.54 for T5 which extends for 1332 km and has very heterogeneous vegetation crossing Pantanal, Amazonia and Cerrado. Biomass dataset in this transect presented high variability while LAI_{dry} presented lower variability (Figure 12). The first four pixels of T5 are located outside of the map of biomass published by Nogueira et al. (2014), and therefore were excluded from the correlation analysis. In the central Amazonia-Cerrado transition, the observed biomass and LAI_{dry} MODIS showed better correlation with higher values (R). The data for all transects were used in the next Section (2.3.3) for comparison with INLAND model simulations.



Figure 12. Longitudinal gradient of the dry season leaf area index (LAI_{dry} – MODIS) and biomass (Nogueira et al., 2014) for the transects 1 to 4 (a) and transect 5 (b).

2.3.3 Longitudinal gradient of biomass and leaf area index in the Amazonia-Cerrado transition – *Comparison between simulations and field observations*

The comparison of simulated results considering the different combinations between nutritional limitation (PR and PG), occurrence of fire (F), interannual climate variability (CV) and average climate (CA) against the dry season LAI (MODIS- LAI_{dry}) and biomass data are presented in this Section.

All transects simulated show overestimated LAI (sum of LAI_{upper} and LAI_{lower}) values and underestimation of biomass values (Appendix, Figures A1 and A2). Although the difference between simulated and observed values are not small, the INLAND model was able to adequately represent the longitudinal gradient for all transects, with higher values to the west and lower LAI and biomass values to the east. The correlations between simulated and observed values of LAI and biomass were higher when considering the interannual climate variability scenario (CV) for most transects, with average values of 0.49 for LAI and 0.71 for biomass (Tables 9 and 10).

Transects **T1 T2 T3 T4 T5** average CA CV CA CV CA CV CA CV CA CV CA CV PC 0.25 0.25 0.47 0.45 0.85 0.91 0.80 0.84 0.39 0.39 0.55 0.57 PR 0.22 0.22 0.43 0.39 0.83 0.89 0.81 0.29 0.42 0.49 0.54 0.46 0.26 0.26 0.55 0.52 0.78 0.84 0.75 0.78 0.53 0.59 0.58 PG 0.60 PC+F0.32 0.14 0.44 0.38 0.71 0.92 0.72 0.84 0.28 0.19 0.50 0.49 PR+F 0.30 0.22 0.39 0.20 0.57 0.78 0.47 0.62 0.25 0.24 0.40 0.41 0.34 0.10 0.50 0.44 0.48 0.79 0.36 0.81 PG+F 0.35 0.35 0.41 0.50 PC+F+CO₂ 0.18 0.17 0.43 0.39 0.72 0.91 0.70 0.83 -0.06 0.06 0.39 0.47 PR+F+CO2 0.38 0.21 0.31 0.33 0.41 0.74 0.69 0.79 -0.07 0.08 0.34 0.43 PG+F+CO₂ 0.40 0.24 0.45 0.37 0.52 0.71 0.60 0.80 0.03 0.20 0.40 0.46 0.29 0.20 0.44 0.39 0.65 0.83 0.66 0.73 0.24 0.29 0.46 0.49 Average

Table 9. Correlation coefficients of dry season LAI simulated by INLAND and remote sensing estimates by MODIS.

	T1		Т	T2 T3		Г3	T4		Т5		Transects	
		_	12				-	average				
	CA	CV	CA	CV	CA	CV	CA	CV	CA	CV	CA	CV
PC	0.88	0.83	0.56	0.53	0.84	0.87	0.79	0.80	0.26	0.30	0.67	0.66
PR	0.87	0.82	0.42	0.39	0.82	0.84	0.78	0.78	0.34	0.41	0.65	0.65
PG	0.87	0.82	0.57	0.54	0.79	0.81	0.71	0.71	0.39	0.44	0.67	0.66
PC+F	0.76	0.80	0.93	0.91	0.90	0.91	0.73	0.71	0.12	0.35	0.69	0.74
PR+F	0.72	0.76	0.92	0.91	0.90	0.92	0.75	0.74	0.12	0.35	0.69	0.74
PG+F	0.71	0.80	0.93	0.92	0.91	0.93	0.64	0.72	0.15	0.39	0.67	0.75
PC+F+CO ₂	0.78	0.80	0.93	0.91	0.84	0.92	0.78	0.73	0.23	0.33	0.71	0.74
PR+F+CO ₂	0.66	0.77	0.90	0.89	0.95	0.85	0.74	0.73	0.35	0.34	0.72	0.72
PG+F+CO ₂	0.63	0.76	0.91	0.91	0.91	0.94	0.54	0.72	0.21	0.34	0.64	0.74
Average	0.76	0.80	0.79	0.77	0.87	0.89	0.72	0.74	0.24	0.36	0.68	0.71

Table 10. Correlation coefficients of biomass simulated by INLAND and field estimates biomass (Nogueira et al., 2014).

The LAI correlation considering only nutritional limitation was higher in PG (0.60) than when using regional phosphorus (PR, 0.46) in the average of the transects (Table 9). Also in the biomass comparison (Table 10), the simulated data showed higher correlation with the observed data in PG (0.66) was in PR (0.65), reinforcing that the use of PG represented the observed data better than PR.

When incorporating the effect of fire (PC+F; PR+F; PG+F), it is possible to observe a decrease in correlations for LAI (Table 9) and an increase in correlations for biomass (Table 10) for all nutritional scenarios. With the fire effect, the best correlations were PG+F with 0.50 for LAI and 0.75 for biomass.

On average for all transects, the combined effect of nutritional limitation, fire and CO₂ for LAI showed higher correlation for PC+F+CO₂ (0.47) than for PR+F+CO₂ (0.43) and for PG+F+CO₂ (0.46) (Table 9). PC+F+CO₂ and PG+F+CO₂ have similar correlations (0.74) for biomass (Table 10). The addition of the transient CO₂ factors to simulations decreased correlations for LAI and biomass, in relation to considering only fire and nutritional limitations. The average correlations for LAI at transects T1 and T2 – northern part of the transition area – were, respectively, 0.20 and 0.39, that are considered low. On the other hand, the average correlations for biomass in these transects were considered high, 0.80 and 0.77 for respectively T1 and T2. This difference suggests possible distortions in LAI_{dry} values, once a high correlation was found for the biomass. It is known that the products of the MODIS sensor in higher rainfall regions are influenced by cloudiness, which affects the accuracy and robustness of the information collected on these regions. The average LAI values used, however, corresponds to months of lower precipitation and cloudiness, which minimizes this potential interference. On the other hand, it is possible that the lower correlations are due to low variations of the simulated LAI values by the INLAND model between the rainy and dry months. The carbon increase in the leaves is controlled by the model's vegetation dynamics module that adjusts the LAI values annually. Possibly, this module may not change in appropriate proportions the LAI in these areas.

The T3 and T4 transects located in the central region of the Amazon-Cerrado transition showed high average correlations between observed and simulated data for both variables, LAI and biomass, when CV were considered (Table 9 and Table 10). The average correlation values for LAI were 0.83 for T3 and 0.73 for T4, and for biomass were 0.89 for T3 and 0.74 for T4, showing high affinity between simulated and observed values. In these regions, that show greater amplitude of the precipitation annual cycle, the model was better suited to simulate variations of LAI, NPP and biomass values. In regions with lower amplitude and variation in annual precipitation, lower average correlations were found. This suggests that the model did not capture the real variation of LAI because it did not seasonally change the rate of photosynthesis

enough to affect the feedback system – similar to that previously discussed for the transect T1 and T2. The T5 transect, located south of the transition, presented low average correlations for CV and CA. This region, characterized by broad vegetation heterogeneity, makes not only the representation through simulation difficult, but also the collection of large-scale data for validation.

In general, the INLAND model simulations showed high and positive correlations, mainly for transects located in the central region of the Amazon-Cerrado transition. This relationship between the observed data and simulated values was stronger when considering the interannual climate variability, especially for biomass, i.e., the use of an average climate decreases the accuracy of the model in simulating the NPP dynamics, biomass, and consequently LAI.

The highest correlation found, however, was for biomass in $PG+F+CO_2$ in the transect T3 where the correlation coefficient was equal to 0.94. This high correlation to biomass was accompanied by a correlation of 0.71 for LAI, the lowest in T3.

The low correlation in PG+F+CO₂ for LAI in T3 may be connected to the fact that the fire effect on the LAI is, in some ways, a compensatory effect in the INLAND model. It happens because the LAI reduction in the arboreal vegetation implies an increase in the LAI for grasses, which means the model will always have high levels of total LAI. These high LAI values do not necessarily imply a high biomass once LAI can increase quickly after the fire occurrence, which may explain the higher correlations for biomass when compared to LAI.

The correlations between the simulated and observed biomass were better than the correlations between simulated and observed LAI. It means that, while the model is
able to capture the Amazon-Cerrado gradient, this robustness is more due to the estimate of biomass than of LAI.

2.3.4 Simulated composition of vegetation

INLAND describes the structure and composition of vegetation communities based on different plant functional types (e.g., tropical broadleaf evergreen trees or C4 grasses). The model has a dynamic vegetation module and, if this module is used, the vegetation type that dominates a particular region can change dynamically to another based on LAI. The model specifies that tropical evergreen forest and tropical deciduous forest ecosystems are characterized by LAI_{upper} above 2.5 m² m⁻². LAI_{upper} values between 0.8 and 2.5 m² m⁻² characterize savannas, and values smaller than 0.8 m² m⁻² characterize grassland ecosystems.

The previous results showed that biomass is simulated more robustly than the LAI. However, this robustness appears to be more quantitative than qualitative, since the model was able to capture the Amazon-Cerrado gradient with higher LAI values in the west and lower values in the east (Figure 12). The LAI estimated by INLAND for the Amazon was between 7.5 and 11 m² m⁻², and 0 - 6 m² m⁻² for Cerrado (Figure 8). These values are overestimated when compared with MODIS observations, these being respectively between 4 and 7 m² m⁻² for Amazonia, and 0 and 3 m² m⁻² for Cerrado (Figure 7).

Although the simulated LAI was overestimated when compared with MODIS observations, it represents well the distribution of vegetation in some conditions with fire and nutritional limitation (Figures 13 and 14). We evaluated the INLAND ability to assign the type of dominant vegetation analyzing 10 years of probability of occurrence.

If the dominant vegetation type in a pixel is identical in greater than 90% of the simulated years (9 of 10), then the simulated vegetation is defined as "very robust" for that pixel; if it occurs in 70 - 90% of the simulated years, it is considered to be "robust". If the dominant vegetation occurred in less than 70% of simulated years, the pixel considered "transitional" vegetation (Figures 13 and 14).

Most of the pixels showed agreement above 90% in vegetation cover classification for CA (Figures 13b, 13d and 13f). A larger number of pixels with values below 70% agreement was observed when considering CV compared to CA. This increase in the heterogeneity of the pixels is due to a higher variation of annual carbon flux induced by the interannual climate variability that directly affects the LAI.

In addition to the overall decrease in robustness, we observed a higher variability in CV compared to CA simulations when we added different effects (P, F or CO₂). Probably, the effect of these factors associated with climate variability acted more strongly on the plants' physiological processes and directly affected the NPP, biomass and LAI. The composition of the vegetation in all nutritional limitation only scenarios for CA simulations resulted in >90% of agreement on the vegetation type for nearly all pixels (Figures 13b, 13d and 13f). CA+PC and CA+PR simulations had the same composition of the vegetation, while CA+PG replaced the deciduous forest by evergreen forest in the central Cerrado region, around 8° S 46° W (Figures 13a, 13c and 13e). Cerrado was better represented in CV+PC, CV+PR and CV+PG than in the same CA combinations (Figures 14a, 14c and 14e). The occurrence of forested areas in the central Cerrado decreased in CV+PC, these being replaced by the savanna vegetation class (Figures 14a, 14c and 14e). However, some areas of perennial forest remained in the central Cerrado in CV+PG, as in CA+PG (Figure 14e). When the effect of fire was added to CA simulations, we can observe an increase in uncertainty on the vegetation cover classification in the Cerrado region. The effect of fire reduced the presence of deciduous forest in central Cerrado biome when compared to PC and PC+CO₂, and the vegetation was replaced by evergreen forest and savanna (Figures 13m, 13o and 13q). In CV simulations, the fire effect results in the replacement of the deciduous and perennial forest by savanna and grasses in all Cerrado central region (Figures 14m, 14o and 14q). This replacement indicated an improvement in the representation of vegetation classes in the Amazon-Cerrado transition.

Few changes in the vegetation composition due to the CO₂ influence were found in the border region when both climate scenarios were compared with the control simulation. However, the CO₂ changed vegetation classes in some locations in the central Cerrado. The major portion of the deciduous forest in CA+PC and CA+PR (Figures 13g and 13i) was transformed in evergreen forest and savanna in CA+PC+CO₂ and CA+PR+CO₂ (Figures 14g, and 14i). The differences between CA+PG+CO₂ and CV+PG+CO₂ are smaller but can be noted in the central Cerrado region.

Finally, we analyzed the effects of the combinations of nutritional limitation, fire and CO_2 on vegetation. The Amazon transition vegetation was better represented in CV and the border of the majority of the savanna class roughly matched the Amazon-Cerrado border set by PROBIO in this simulation (Figure 14x). This combination also results in savanna and grasses dominating the central Cerrado. Althought INLAND model overestimated LAI values, this model can represent well the distribution and classification of the Amazon-Cerrado transition under the effects of nutritional limitation, fire and CO_2 when climate variability is considered. This improvement can be confirmed not only by visual analysis of the vegetation composition, but also through improvements in the correlations when considering CV.

Although the model has performed well in representing the composition of the vegetation in all cases, no changes were found for the southern region of Mato Grosso state. There is one small area of evergreen forest, suggesting that this region may have soil physical and chemical properties different from those used in the model. In the northern Amazon, INLAND could represent with >90% robustness the Cerrado areas in all simulations.



Figure 13. Results of the vegetation composition when incorporated the effects of nutritional limitation, fire and CO₂ in CA and simulated pattern of dominant vegetation.



Figure 14. Results of the vegetation composition when incorporated the effects of nutritional limitation, fire and CO₂ in CV and simulated pattern of dominant vegetation.

2.4 CONCLUSIONS

Results from the model simulations indicate that, in INLAND, climate variability and interannual disturbances such as fire have a major impact on the vegetation structure. Processes of competition between trees and grasses for resources are intensified, changing the vegetation cover with little variation in carbon storage.

The experiments conducted to evaluate the effect of nutritional limitation on vegetation in the Amazon-Cerrado transition revealed that phosphorus is more limiting in the Amazon than in the Cerrado, highlighting the importance of incorporating P dynamics in DGVMs, since tropical forests have a determinant role in the global carbon balance. In the Cerrado, on the other hand, the effect of nutritional limitation is smaller but not insignificant, since our correlations showed an improvement in correlation between predicted and observed data for LAI and biomass in the presence of phosphorus limitations.

In the Cerrado region, the effect of fire is much more intense than the effects of the nutritional limitation, so that in the transition region fire can be considered the main agent of change in the vegetation. Fire effects on the structure and dynamics of vegetation are directly linked to their intensity and frequency, which commonly results from a more humid or drier climate. When evaluated in combination with CO₂, the effects of nutritional restriction and fire showed fewer changes in land cover, so its addition brought complexity and nonlinearity to the results.

Although our results have identified a better representation of biomass by the INLAND model compared to LAI, significant improvements in the representation of the Amazon-Cerrado border were found when the effects of phosphorus and fire were considered. The addition of nutritional limitation, fire, and subsequently CO₂, gradually improved representation of the

Amazon-Cerrado LAI gradient, causing the model to simulate the current position of biomes. The best combination for representing the Amazon-Cerrado border was the $CV+PG+F+CO_2$ scenario.

3. GENERAL CONCLUSIONS

In general, we conclude that climate, fire and soil phosphorus have an important role in the dynamics of vegetation along the Amazon-Cerrado transition, changing the net primary productivity, biomass and leaf area index. Although these factors are important, and of great interest for community scientific, the lack of observed data in transition regions is still a major limitation for the improvement of the dynamic global vegetation models. In this work we had difficulty obtaining observed data for physiological parameters such as soil P and V_{max} , for the Amazonia-Cerrado transition, because most of the available observed data for physiological parameters refers to a particular type of vegetation or predominant species, not representing the transition area.

Given this limitation, we use a pre-established relationship between V_{max} and total soil phosphorus total (P_{total}) to estimate V_{max} , remote sensing data to obtain LAI and a biomass map. However, finding soil phosphorus total measurements was also a challenge. Most soil phosphorus data does not follow a common protocol, ie, there are measurements of the available phosphorus in the soil obtained from different extractors, making it difficult to compare and use of this data together. In addition, most of the information about the soil phosphorus content is not accompanied by data as texture, clay content, soil type, which makes it even more complex the use of this information. To solve part of this problem and homogenise the soil data measurements used, we used a mathematical model to estimate the soil phosphorus total from phosphorus regional map in the Amazon-Cerrado transition incorporating phosphorus data collected in the Mato Grosso state and the allowing a numerical experiment with the INLAND model.

From the numerical experiment, it is possible to conclude that the use of climate interannual variability resulted in a statistically significant decrease of carbon increment and improved the correlation between observed and simulated LAI and biomass. Fire showed a statistically significant influence only in the Amazon-Cerrado border, not affecting Central Amazonia. The effects of nutritional limitation were more important in the Amazon than in the Cerrado biome.

When used in combination, however, these factors improved the spatial representation of vegetation classes in the model. These results showed that the INLAND model can be a powerful tool in assessing possible changes in vegetation under scenarios of future climate change and emphasize the importance of a greater number of observed data. To further study the nutritional limitation in transition areas, it is recommended to use Hedley fractionation data obtained for total soil phosphorus, because this method also allows the extraction of more varied forms of available phosphorus in the soil, which can help us to quantify how much of the total phosphorus acts in the nutritional limitation of photosynthesis.

Suggestions for future implementations in INLAND include: a) modifications in the INLAND phenology module improve the representation of the seasonal LAI; b) substitution of the prescribed C:N ratio existing the model by a dynamic soil N cycle, since the phosphorus photosynthesis limitation depends on the nitrogen content in the leaves; and finally c) implementation of a dynamic phosphorus cycle and nonlinear interactions between the phosphorus present in the leaves and soil.

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5. APPENDIX A

Plot	Location		PResin	Ptotal	Clay
ID	Latitude	Longitude	mg kg ⁻¹	mg kg ⁻¹	%
'RIO-12'	8.5	-61.5	3.42	178.96	9.50
'ELD-12'	6.5	-61.5	5.34	173.59	20.13
'SCR-01'	1.5	-67.5	2.36	65.85	6.88
'TIP-05'	-0.5	-76.5	9.01	437.32	37.30
'JRI-01'	-0.5	-52.5	1.30	189.13	80.73
'JAS-02'	-1.5	-77.5	8.79	423.65	29.13
'CAX-01'	-1.5	-51.5	5.17	115.18	41.77
'MBO-01'	-1.5	-48.5	3.14	101.38	11.49
'BNT-04'	-2.5	-60.5	4.82	68.67	57.68
'TAP-04'	-2.5	-54.5	4.50	192.34	89.25
'ALP-12'	-3.5	-73.5	7.37	86.62	13.96
'SUC-02'	-3.5	-72.5	4.06	349.81	37.21
'AGP-01'	-3.5	-70.5	2.99	303.28	42.61
'ZAR-03'	-3.5	-69.5	4.81	177.19	31.11
'TAP-123'	-3.5	-54.5	1.65	78.89	66.10
'ZAR-04'	-4.5	-69.5	7.48	54.15	18.26
'JUR-01'	-8.5	-72.5	10.55	331.93	36.62
'RST-01'	-9.5	-72.5	8.29	240.19	25.40
'ALF-01'	-9.5	-55.5	3.51	118.14	11.43
'DOI-01'	-10.5	-68.5	7.18	203.19	19.09
'SIN-01'	-11.5	-55.5	2.58	61.25	9.80
'TAM-01'	-12.5	-69.5	5.85	343.51	37.75
'CUZ-03'	-12.5	-68.5	11.45		42.46
'CRP-01'	-14.5	-61.5	21.82		18.05
'HCC-21'	-14.5	-60.5	7.34	289.76	25.60

Table A1. Phosphorus samples in P_{resin} and clay fraction to different plots in the Brazilian Amazon from the database Quesada et al. (2010).



Figure A1. LAI observed and simulated for the five transects in the Amazon-Cerrado transition.

