Temporal rainfall variability as inductor of the phenology of Brazilian semiarid species

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Abstract

Usually, in the Brazilian semiarid region where dry and humid periods are well determined, phenological events are mainly affected by seasonal rainfall and availability of water in the soil. The studies in these regions have importance especially to understand the dynamics of dry forests. Thus, the objective of this work was to characterize the phenophases of trees P. pyramidalis, C. blachetianus, A. pyrifolium and J. mollissima, and their relationship with the temporal distribution of rainfall, which occurs in the form of pulses. Furthermore, the effect of goat grazing was evaluated. The experimental areas presented different caprine stockings, where first had ten and the second five. The third stocking remained without goats during the experimental period. For the phenological follow-up, 50 tree individuals of each species were selected, tagged and monitored weekly, at Experimental Station, in the municipality of São João do Cariri-PB. In order to quantify the phenological events, the percentage of Fournier intensity was used, registering the presence and absence of the phenophases: sprouting, flowering, fruiting and senescence, whose data were related to the water pulses and inter-pulses. The phenological events presented annual patterns, with interannual variations in intensity. We observed that the phenophases occur in synchrony with the temporal distribution of the rainfall pulses in the area of its occurrence. As the dry season progresses, reduction in sprouting, flowering and fruiting occurs when the inter-pulses of rainfall was intensified. The presence of goats did not influence the phenological development of the species.

Keywords: Dry forests; Fournier intensity; Goats; Phenophases; Water pulses and inter-pulses.
Abbreviations: FI_Fournier intensity.

Introduction

Phenology helps in understanding the dynamics of forest ecosystems and is essential for the ecological and evolutionary study of the tropics. Thus, it is necessary to know the periodicity of vegetative and reproductive events to provide subsidies that aid in the complex dynamics of ecosystems (Braga et al., 2019). Thus, phenological studies are important to evaluate the periods that the plant presents flowers and fruits, as well as other phenophases, and to provide information about the life cycle of these plants, being a crucial tool in the knowledge of the dynamics of these organisms (Paz and Souza et al., 2018).

The Brazilian semiarid region is characterized by a high irregularity of the rainfall distribution in space and time. It has two distinct seasons, the rainy season, corresponding to a short period with rainfall, erratic and of very variable intensity; and the dry season, in general with critical period of prolonged drought, when the Caatinga species express their most striking physiognomy, the deciduous. In these conditions, the plant species present dynamics of growth and development in consonance with the rainfall events (Lima et al., 2018).

The Caatinga provided a scenario for the evolution and irradiation of unique plant strains that contrast with tropical rainforest humid lineage as the plants in this dry forest have developed exceptional adaptations to cope with a “hostile” environment with prolonged, irregular rains and droughts. Several studies have shown that precipitation controls the phenology of many Caatinga species, although other species may start their phenophases independently of the occurrence of rain (Silva et al., 2018).

In the Caatinga, there are woody species, in which the leaf abscission occurs at the end of the rainy season, while other species maintain their leaves during the dry period, resulting in different tree species over time and in space (Silva et al., 2017). It should also be considered that precipitation events are never uniformly spaced in time, and that the probability of occurrence of these events increases during the dry season. It has also been suggested that dry forests provide goods and services significantly different from those of
humid forests, requiring different approaches to management and conservation (Brand, 2017). Typically, rainfall events cause concise pulses of soil water content from infiltration due to the small effective depth of most soils in the region. It should be considered that although precipitation events are never uniformly spaced in time, the probability of occurrence of these events increases during the rainy season (Noy-Meir, 1973).

In view of the complexity of the phenological events in the Brazilian semi-arid regions, the objective of the present study was to characterize the phenophases of *Pandanus pyramidalis* Balf.f., *Croton blachetianus* Baill, *Aspidosperma pyrifolium* Mart. & Zucc. and *Jatropha mollissima* (Pohl) Baill, endemic to the Caatinga. These species have multiple uses, and their relationship with the temporal distribution of rain pulses is important. Furthermore, evaluation of the effect of the goat on grazing is carried out as a significant factor.

**Results and discussion**

**Sprouting (emergence of leaves)**

In the first year of evaluation (2007) different intensities were observed for sprouting in the studied species, in which *P. pyramidalis* presented higher Fournier intensity than the others (Fig 1). The sprouting period of *P. pyramidalis* was occurred between March and July, varying between years, with rapid leaf emission in response to the precipitation pulses, peaked a few days after the rains. Souza et al. (2014) and Parente et al. (2012) identified that rainfall pulses are sufficient to trigger phenological events in this species. Thus, sprouting of *P. pyramidalis* can be classified as a rapid and intense precipitation-dependent event in sufficient quantity to rehydrate the plant.

In the individuals of *C. blachetianus*, the first rains represented the necessary stimulus to trigger the leaf emission that occurred during the months of February to July, with differences of a few days between the study areas (Figure 1). In 2007, the individuals of *C. blachetianus* re-grew after sporadic rains. In the years 2008, 2010 and 2011, no such behavior was observed, corroborating with the hypothesis that probably the leaf buds of this species enter a state of dormancy.

In contrast to our results, Amorim et al. (2009) observed the absence of dormancy of leaf blossoms in *C. blachetianus* plants in Brazilian semi-arid region. They indicated that the resumption of foliar emission is complex and involves many other factors other than response to water availability. This demonstrates how much still needs to be studied on the phenological patterns of Caatinga species.

The sprouting pattern of *A. pyrifolium* presented continuous during the months of February to July, varying according to the years, beginning just before the rainy season (Fig 1). *A. pyrifolium* initiated the leaf loss from July, remaining without leaves for a long period. Another characteristic of this species that was also noticed in *C. blachetianus* is the response to rain events, waiting for more intense rains to restart the leaf emission.

Amorim et al. (2009), worked with another species of the genus *Aspidosperma*, and reported that both sprouting and leaf fall presented a continuous pattern, with the number of leaves being reduced in relation to sprouting. Parente et al. (2012) studied the same species and identified that the precipitation interfered with phenological behavior, triggering the sprouting period soon after the first rainfall events, in which *A. pyrifolium* was more persistent in maintaining the leaves with the prolongation of the dry season. The higher leaf persistence of *A. pyrifolium* individuals may be associated with the amount of nitrogen translocated to the new leaves as observed by Castro et al. (2013) when studying the phenology of *Aspidosperma* species.

The species *J. mollissima* showed sprouting between the months of March and June, reaching the peak intensity of Fournier in April 2010 (Figure 1). The species also responded positively to rainfall, presenting a short period of leaf maintenance, being sensitive to the water deficit. This is corroborating with Fernandes et al. (2013) that associated the formation of new leaves to the rainy season and the leaf fall to dry season.

On the other hand, Neves et al. (2010) studied three species of *Jatropha* in distinct Caatinga environments and observed continuous phenological behavior of the individuals due to the capacity of water storage in the stem. This characteristic allows the occurrence of phenological events even in the absence of rainfall. Even *C. blachetianus* and *A. pyrifolium*, are consumed in the dry season, the presence of goats had no effect on the phenology of the individuals. This disagrees with the reports by Parente et al. (2012) who studied the influence of grazing on the phenology of four species in Caatinga area and verified a reduction in total leaves, fruits and flowers in the grazed pickets.

However, this divergence can be explained by the methodological form, in which the phenological data were analyzed. Parente et al. (2012) studied results obtained through weighted averages of visual observations within a period of two years encompassing longer period of phenological evaluation of the species. They observed that the action of grazing goats in the experimental areas did not interfere in the phenological development of the species.

**Leaf senescence**

The recorded senescence period for *P. pyramidalis* varied between May and September, according to the interval between the pulses of precipitation (Fig 2). The phenological behavior was similar to that reported by Souza et al. (2014) and Parente et al. (2012) in study of Caatinga species. *P. pyramidalis* begins the foliar fall in a concentrated way at the end of the dry season. The tendency of the trees of dry environments, where the plants presented their phenophases was mainly regulated by the precipitation.

During the years of 2007 and 2008, we observed that *P. pyramidalis* remained without leaves in a period between beginning of August until the month of January, coinciding with the drought season, when the rain begin to become scarce, and soil moisture causes the plant to stabilize the sprouting and initiate foliar abscission. The sensitivity of *P. pyramidalis* to variations in rainfall can be explained by the fact that this species has a dense stem. So, that the possibilities of water reserves become limited, resulting in greater sensitivity to soil water changes and to the water deficit, as measured by interpulse predominates over the precipitation pulses. According to Nogueira et al. (2013) and Santos et al. (2013), dense wood species are sensitive to
water stress. They rapidly lose leaves as a way to reduce moisture loss.

Figure 2 showed that foliar abscission in *C. blachetianus* individuals are occurred more continuously over the years, from February to November, when precipitation interpulse became predominant, causing a reduction in soil water content. The period in which *C. blachetianus* remains without leaves is comprised between July and August, corroborating Amorim et al. (2009), who cited this species as being regular in leaf fall.

Leaf fall in the species *A. pyrifolium* is observed during the months of May to September, being intensified in the most critical period of soil water scarcity, as recorded in the other species (Figure 2). The senescence period of the species *Aspidosperma* can extend this phenophase until October and present in their constitution, reduced amounts of nutrients, mainly nitrogen and carbon, recruited for the following phenophases of flowering and later fruiting.

The leaf fall pattern of *J. mollissima* occurred between April and July, remaining without leaves for three months. The intensity of leaf abscission during the years 2010 and 2011 was reduced, probably due to the distribution of rainfall (Figure 2). Amorim et al. (2009) correlated the senescence pattern of *J. mollissima* with *C. blachetianus*, classifying them as similar, although the leafless period of *J. mollissima* is between two and three months on average, smaller than that of *C. blachetianus*.

Fernandes et al. (2013), worked with *J. curcas*, observed a behavior similar to that reported for *J. mollissima* in the years 2010 and 2011, during the rainy season, when there was a significant reduction in leaf abscission. According to Maes et al. (2009), the leaf abscission of *J. mollissima* guarantees the efficient use of water in the dry season, as well as in the other Caatinga species, beginning at the end of the rainy season and during the dry season.

**Flowering**

In the population of *P. pyramidalis*, the flowering period was recorded between March and June (Figure 3) when the leaves were fully expanded. According to Souza et al. (2014) the duration of flowering of *P. pyramidalis* can occur in the period of six months, varying according to the environmental conditions, corroborating with the results obtained in this study.

Flowering peaks in *P. pyramidalis* individuals were observed in May 2008 and 2010 and April 2011. In 2007, even the evaluation period occurred at the end of the rains, the peak was observed in the month of July. Although shorter than the other phenophases, flowering peaks recorded in individuals of *P. pyramidalis* may represent a strategy to concentrate the presence of seasonal pollinators (Van Schaik et al., 1993). The timing occurrence of reproductive phenophases, especially flowering, is crucial because it influences pollination success and thus the rate of seed production (Talora and Morelatto, 2000).

Flowering of *C. blachetianus* was occurred in the months of January to June, with peaks in April (2008), March (2010) and February (2011), coinciding with half of the rainy season (Figure 3). In this study, we observed that the presence of goat grazing did not influence the occurrence of these phenophases, since according to Leal et al. (2003), goats look for young seedlings more than flowers and fruits.

*A. pyrifolium* showed the highest flowering period among the studied species. The flowering was recorded between February and December, with peaks in March (2008) and February (2010). In 2011, flowering showed a greater uniformity in the distribution of this phenophase (Figure 3).

The flowering pattern recorded for *J. mollissima* was annual, lasting between February and June, with peaks in March, February and May, for the years 2008, 2010 and 2011, respectively, varying according to the period and duration of the rainy season (Figure 3). The performance of climatic variables on flowering and other phenophases of *J. mollissima* is reported by Neves et al. (2010), Jong.schema et al. (2007) and Santos et al. (2005) who observed absence of flowers, fruits, leaf fall and sprouting according to the availability of water in the soil.

As observed by Torres and Galetto (2011) and Lesica and Kittelson (2010), in dry areas, flowering tends to occur in the rainy season. In the Caatinga, Amorim et al. (2009), Parente et al. (2012) and Souza et al. (2014) revealed the tendency of the woody species to bloom in the middle or end of the rainy season.

Amorim et al. (2009) observed that flowering in the species *C. blachetianus* and *J. mollissima* occurs along the leaf flow phenophase. Machado et al. (1997) reported that flowering can also be observed even before leaf formation. In order to understand the variability of flowering patterns of species action of climatic factors, genetic and intrinsic factors is vital, which highlights the importance of further studies.

**Fruiting**

Fruiting in the individuals of *P. pyramidalis* was observed between the months of May and July, in the middle of the rainy period, with peaks in the months of July (2007, 2008 and 2010) and May during the year of 2011 (Fig 4). Souza et al. (2014) reported that the fruiting period in *P. pyramidalis* individuals may extend up to eight months. However, in the evaluated individuals the fruiting period (three months) began in the middle of the rainy season of Caatinga, Brazilian semiarid region.

The fruiting period is directly related to the conditions that influence the success of the dispersion and establishment. The flowering season in seasonal forests is often related to the type of fruit and the dispersal syndrome. For example, *P. pyramidalis* (which is nut), usually fruit in the middle or end of the rainy season (Schwartz, 2013).

*C. blachetianus* presented fruiting period in the months of February to July, in the years 2008 and 2010, where the peaks reached 60% of intensity (Fig 4). It is important to emphasize that there was no difference in the areas as the duration of fruiting peaks, indicating that the presence of goats does not influence the fruitful development of the species, until the stocking of up to ten animals.

According to Fig 4, the fruiting period of *A. pyrifolium* comprised of April to September. In the year 2008, a short but well defined fruiting peak was identified different from the following years, where a continuity of the event is observed, in area with ten animals. Area I, presented a slightly superior performance compared to other areas, contrary to what was reported by several authors. The goats
Fig 1. FI (%) for sprouting of Caatinga tree species, with different goat stockings from June 2007 to September 2011.
Fig 2. Fi (%) for leaf senescence of Caatinga tree species, with different goat stockings from June 2007 to September 2011.
Fig 3. FI (%) for flowering of Caatinga tree species, with different goat stockings from June 2007 to September 2011.
**Fig 4.** Fi (%) for fruiting of Caatinga tree species, with different goat stockings from June 2007 to September 2011.
had no influence on the reproductive phenology of this species, among other reasons, to the chemical and anatomical constitution of the fruits. The occurrence of fruiting during the whole year suggests that environments with absence of dry periods offer little restrictive conditions for the continuous development and maturation of fruits (Bauer et al., 2012). In species such as *A. pyrifolium*, fruiting and dispersal in the rainy season is favored by keeping the fruits more attractive to the dispersers for longer periods, favoring greater zoological dispersion (Belo et al., 2013, Lenza and Klink, 2006; Batalha and Mantovanni, 2000).

*J. mollissima* presents the fruiting between March and June with well-defined peaks annually (Fig 4). Santos et al. (2010) studied the reproductive phenology of the *J. curcas* observed a peak of flowering and fruiting during the rainy season, began in April and extending until August, with peaks during the month of June and July, similar to that reported by this work.

The species *P. pyramidalis*, *C. blachetianus*, *A. pyrifolium* and *J. mollissima* present low dispersion in fruiting events, with average CPD below 40%. This data indicates that the occurrence of the phenophases was occurred for short periods, corroborating with the length of the mean vector. It shows high concentration of flowering at certain times of the year.

The results of the average vector length with concentration around the mean dates high for the years 2007 and 2008, indicates seasonality. Santos et al. (2010) also reported that the vegetative and reproductive events of Caatinga species are mostly seasonal, concentrating their development more intensely in the rainy season.

### Materials and methods

The field study was conducted on Experimental Station of the Federal University of Paraíba, located in the municipality of São João do Cariri, Paraíba State (PB), Brazil, at coordinates -7º 23’ 30” latitude and -36º 31’ 59,” longitude. The predominant relief in the area is mildly undulated with an altitude of approximately 458 m in relation to the sea level.

The region is characterized by having a hot and dry climate; according to the Köppen (1948) classification. It is a semiarid type, BSh subtype, with an average annual precipitation of 529 mm, maximum and minimum temperatures of 31 and 17.2 °C, respectively, with small variations over the year, and relative air humidity around 77%.

Over the course of the experiment, daily records of precipitation were collected at Climatological Station, in São João do Cariri/PB.

The soils of the study area are mostly shallow, relatively plain to mildly undulate. They are on a crystalline geological structure where the most present types are: Chromic vertic luvisol – tc (Noncalic brown vertic), Chromic orthic vertisol – vc (Vertisol) and Lithic entisol – rl (Lithic soil). There is also, in small proportions, Eutric Cambisol – cx (Cambisol), Eutric Fluvisol – ru (Alluvial soil), Haplic planosol – sx (Planosol sco) and rocky outcrops with evidence of Lithic entisol (Chaves et al., 2002).

The experimental area comprised of 9.6 ha, divided into three paddocks of 3.2 ha each, with different capacities: Area I (10 goats); area II (five goats); and area III (without the presence of goats). Goats were adult, male and without defined racial pattern, received daily concentrated and mineral supplementation in amount of 200 g per animal.

### Plant materials

To select individuals of *P. pyramidalis*, *C. blachetianus*, *A. pyrifolium* and *J. mollissima*, the sampling method was the quadrat (an adaptation of the ‘plots’ method): ten quadrants (approximately 10 × 10 m) within the experimental area were selected so as to represent the entire Caatinga area of the Experimental Station. The criterion for inclusion of individuals was marked as reference spot in every quadrant and selecting the five closest individuals in the north, south, east and west directions. For each species, 50 trees of similar size, having a trunk with minimum diameter at breast height greater than or equal to 20 cm were selected. Plants were identified with aluminum plates and non-woven fabric (TNT) ribbons to be better visualized.

The observations were performed in weekly intervals, recording the presence and absence of the sprouting, leaf senescence, flowering and fruiting phenophases. These phases were defined as follows: sprouting (emergence of leaves) – initially characterized by the presence of leaf primordia, usually of light-green or reddish color, ending when leaves acquired a dark green color; senescence – period when the leaves change their color from dark green to brown and the spontaneous leaf abscission starts, forming empty spaces (flaws) at the crown or branches; flowering – period when the tree is at full bloom; fruiting – starts from the moment it is possible to visualize the small fruits after the flowers are fertilized and ends with dispersion of the seeds.

### Fournier’s intensity percentage

It proposed by Fournier (1974), whereas the values obtained in the field with a five-category (0 to 4), semi-quantitative interval scale with an interval of 25% between each category allowed us to estimate the percentage of the phenophase intensity in each individual. Weekly, the intensity values obtained for all individuals of the species were summed and divided by the highest maximum possible value (number of individuals multiplied by four). The result, corresponding to a ratio was then multiplied by 100 to be converted into percentage terms.

### Statistical analysis

The phenological data of intensity of each phenophase (sprouting, senescence, flowering and fruiting) were related to the distribution of rain pulses and through Spearman’s correlation coefficient (Zar, 1996), using the number of individuals under each phenophase and the monthly climate data.

### Conclusions

The annual variation in the occurrence of phenological events recorded in the analyzed species is in accordance with the temporal distribution of rainfall. The intensity of the phenophases is associated to the frequency of the precipitation pulses. The species *P. pyramidalis* is the most sensitive to the variability of precipitation among the species studied. The
presence of goats under grazing does not influence the phenological development of the species analyzed.

References


