Cacti community structure in a tropical Mexican dry forest under chronic disturbance

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Abstract: When agricultural lands in formerly dry forest are abandoned, chronic disturbance by livestock grazing and wood extraction may set back successional processes precluding older dry forest structure from be reached. We compare cacti density, richness, diversity and evenness in twelve 50 × 50 m plots: six plots in secondary dry forest and six plots in older conserved dry forest at Sierra de Huautla, Central Mexico. Cacti were identified, the life-form of each species was recorded and size was measured to assign individuals into four life-stage classes. Nine cacti species were registered across both forest types. In this dry forest, richness, diversity and evenness of cacti community increased in sites under chronic disturbance, whereas density was not significantly affected. Size structure at community level in the conserved forest showed an increase of individuals with size, whereas in the secondary forest, size structure showed an uneven distribution of individuals, as in populations that regenerate by pulses. In the secondary forest, there was a lower proportion of adults and they were in average, smaller than those in the conserved forest. Chronic disturbance may eventually change cacti community structure, affecting animals that depend on large arboreal cacti for food, perches and/or refuge; therefore, maintenance of biodiversity will be at stake.

Key words: cattle ranching, Opuntia spp., Pachycereus grandis, restoration ecology, secondary dry forest, Sierra de Huautla.

The Cactaceae family includes more than 2,000 species of many life forms, from smaller prostrate species to large arboreal ones (Barthlott and Hunt, 1993). Mexico has around 63 genera and 913 taxa (669 species and 244 subspecies); a high percentage of those species (35%) are considered threatened by the destruction of their habitat (Hernández and Godínez, 1994; Hunt, 1999; Guzmán et al., 2003). Worldwide analysis has shown that Mexico is the most important country to preserve cacti diversity in different ecosystems as deserts, xerophitic shrub lands and...
tropical dry forests (Bravo-Hollis, 1978; Bravo-Hollis and Sánchez-Mejorada, 1991a,b; Anderson, 2001; Ortega-Baes and Godínez-Álvarez, 2006). In tropical dry forests, Cactaceae is the third most important family by number of species (Trejo-Vázquez, 1998); however, this family is seldom evaluated in this ecosystem, which is indeed the third one with the highest forest cover loss worldwide (Hansen et al., 2010). Given that tropical dry forests are dominated by wind-dispersed species, cacti represent an important source of fleshy fruits for birds and bats during the dry season (Naranjo et al., 2003; Reyes-Agüero et al., 2006) when most of the tree species do not produce fruits (De León-Ibarra, 2005; Martínez-Garza et al., 2011). Therefore, the conservation of this family is crucial to maintain ecosystem processes, as trophic chains.

Chronosequences studies in tropical dry forests have shown that a forest structure similar to that of mature forest may be reached after more than 30 years of abandonment of agricultural land (Kennard, 2002; Ruiz et al., 2005). However, young secondary forests originated from abandoned agricultural land where successional processes took place (Brown and Lugo, 1990; Finegan, 1996) may experience chronic disturbance; for example, in many areas with secondary forest, cattle graze and browse herbs and grasses at the understory and also at the lower foliage of trees (Stern et al., 2002). Additionally, disturbance due to extraction of wood for timber, firewood or fence posts is frequent (Linares-Palomino and Ponce, 2005). This kind of forest, with different ages and degrees of disturbance, now cover more area than conserved older forest in the tropics (Chazdon et al., 2007). After lands are no longer used for cultivation, chronic anthropogenic disturbance may continuously set back successional processes, precluding mature forest structure to be reached and threatening cactus species inhabiting those areas.

Disturbance may have positive or negative effects on populations of cacti. Some studies have shown that land use transformation and cattle ranching are the main factors affecting negatively the population dynamics of cactus species (e.g. del Castillo, 1987; Bowers, 1997; Martorell and Peters, 2009; Flores et al., 2010). Other studies have revealed that moderate levels of human activities may increase densities of some globose species (Martorell and Peters, 2005; Portilla-Alonso and Martorell, 2011). Demography of cactus species have been frequently studied (reviewed in Godínez-Álvarez et al., 2003) but its community structure and the effect of disturbance on it has not been evaluated. We may expect a differential effect of disturbance in the structure of a community depending from dominant life-form. For example, diversity, richness, and evenness of herbs and grasses increase under extensive cattle ranching in pasturelands (Nai-Bregaglio et al., 2002; Floyd et al., 2003; Hayes and Holl, 2003), whereas livestock presence affects negatively tree richness and diversity in the dry forests (Roth, 1999; González-Iturbe et al., 2002; Stern et al., 2002; VanderWerf, 2004). On the other hand, selective logging may not affect tree evenness in the dry forest (Monroy-Ortiz, 1997), but livestock presence does (Sánchez-Velasquez et al., 2002). However, it is not known whether the effect of disturbance on cactus communities inhabiting the dry forest will follow patterns found in other ecosystems under chronic disturbance.

Population size structure is affected by recruitment and mortality rates. For cactus species, population size structure is frequently an uneven distribution of individuals in the different size classes (see Appendix 1 in Godínez-Álvarez et al., 2003). The uneven population size structure is associated to events of recruitment in pulses favoured by the occasional coincidence of some suitable biotic and abiotic conditions, as rain, temperature, low herbivory and/or presence of nurse plants (Godínez-Álvarez et al., 2003). A couple of studies in the dry forest have revealed that cacti show an uneven size distribution (Contreras and Valverde 2002; Méndez et al., 2004). Other studies, also in the dry forest, have shown a decreasing number of cactus individuals with size (Ferrer-Cervantes et al., 2012; Valencia-Díaz et al., 2012) which is related to constant regeneration and lower mortality of adults (Godínez-Álvarez et al., 2003). Irrespective of population size structure, neutral effect of disturbance on size structure of cactus species have been reported (e.g. Ureta and Martorell, 2009). However, given that cattle may step on smaller life-stage classes (del Castillo, 1987), regeneration of cacti in arid environments may be reduced and thus, individuals reaching adult stage may be fewer compared to habitats experienced undisturbed succession. Therefore, chronic anthropogenic disturbance in older secondary forest may result in an overall size structure similar to the one in the conserved forest but with a lower proportion of individuals from older life-stages.

We investigated cacti community structure in two localities of the tropical dry forest of Sierra de Huautla, Morelos, central Mexico, in fragments of older conserved dry forest and in successional areas more than 30 years old which are frequently disturbed by livestock activities and selective extraction of wood (called from now on, secondary forest). We tested the following hypothesis: due to chronic disturbance, (1) density, diversity, richness, and evenness of cacti community differ in older conserved compared to secondary forest, and (2) the cacti community in the dry forest shows an uneven size structure with a lower proportion of adults in the secondary forest compared to the older forest.

Methods

Study area. This study was carried out in lands close to the localities of El Limón de Cuauchichinola (1,220 m of altitude, called from now on El Limón) and Quilamula (1,070 m of alti-
titude), both located within the Sierra de Huautla Biosphere Reserve, Morelos, central Mexico. The Reserve (18° 20' 10" - 18° 34' 20" N and 98° 51' 20" - 98° 08' 15" W) comprises 59,030 ha where the main vegetation type is tropical dry forest. Mean annual temperature is 24.5 °C and average annual rainfall (1980-2001) amounts to 840 mm (CONAGUA, Gerencia Regional Balsas, unpublished data), ~90% falling between late May and October. During the dry season (November to April), most of the trees lose their leaves. Soils are shallow (< 30 cm in depth), with sandy-loam texture (CONANP, 2005). In the Reserve, the most important plant families by decreasing number of species are Fabaceae, Poaceae, Asteraceae, Burseraceae, and Cactaceae (CONANP, 2005). More than 70% of species and genera of cacti reported for Morelos has been recorded in the Reserve: 12 genera and 28 species (Martínez-Alvarado, 1985; CONANP, 2005).

Most of the trees in this forest have compound leaves and canopies. The most common canopy trees in the Reserve (18° 20' 10" - 98° 08' 15" W) comprises > 70% of species and genera of cacti reported for Morelos has been recorded in the Reserve: 12 genera and 28 species (Martínez-Alvarado, 1985; CONANP, 2005). Most of the trees in this forest have compound leaves and canopies. The most common canopy trees in the Reserve (18° 20' 10" - 98° 08' 15" W) comprises > 70% of species and genera of cacti reported for Morelos has been recorded in the Reserve: 12 genera and 28 species (Martínez-Alvarado, 1985; CONANP, 2005). Most of the trees in this forest have compound leaves and canopies. The most common canopy trees in the Reserve (18° 20' 10" - 98° 08' 15" W) comprises > 70% of species and genera of cacti reported for Morelos has been recorded in the Reserve: 12 genera and 28 species (Martínez-Alvarado, 1985; CONANP, 2005).

The landscape in the Reserve is a mosaic of fragments of old and young tropical dry forest in different degrees of conservation, surrounded by agricultural lands and small towns. Two main forest types are recognized in the localities studied here: secondary forest and conserved forest. The secondary forest resulted from large parts of forest cleared more than 40 years ago (~1970), which were used for maize cultivation for ca. 6 years, and later abandoned. The secondary forest has an average tree density (≥ 5 cm diameter at breast height: 1.3 m) of 264 ind./ha from 14 species, from which Acacia cocciaefalacia (Fabaceae), Ipomoea pauciflora (Convolvulaceae), Acacia farnesiana (Fabaceae), and Mimosa benthamii (Fabaceae) are dominant (Martínez-Garza et al., 2011). These forests are chronically disturbed by anthropogenic activities: during the rainy season ca. 600 heads of livestock feed in these secondary forests (~7 heads/ha); these heads of livestock are brought from neighbouring towns. Accordingly to land owners and current research (B. Maldonado, unpublished data), cattle are maintained in farms during the dry season but goats, pigs and horses graze in the secondary forest throughout the year.

What we call conserved forest corresponds to older growth forest never deforested where cattle from neighbouring towns do not graze, although cattle from local owners occasionally do; nonetheless, cattle is never found in densities higher than 1 head/ha (G. Pacheco, local farmer, pers. comm.). In conserved forests, wood extraction is seldom practiced since the preferred tree species are not frequent (B. Maldonado, unpublished data). The most common canopy trees in the conserved forest are Conzattia multiflora, Lysiloma acapulcense, L. divaricata (Fabaceae), several species of Bursera (Burseraceae) and Ceiba (Bombacaceae) (CONANP, 2005). In this conserved forest, average tree density is 905 ind./ha (unpublished data). Native herbivores in both forest types are deer (Odocoileus virginianus) and rodents (Lymnis sp., Baiomys sp., Sigmodon sp.; D. Valenzuela, unpublished data).

**Sampling protocol.** In January 2006, 12 plots of 50 × 50 m were selected in both forest types (secondary and conserved forest) in El Limón (N = 6) and Quilamula (N = 6) for a total sampling area of 3 ha. Within each locality, we selected three plots in each forest type with similar slopes (<30%) and soil characteristics (J. Campo, unpublished data). These plots are part of a large project of restoration ecology which aims to favour the coexistence of biodiversity and the economic activities of the region using enrichment plantings and exclusion of disturbance (Martínez-Garza et al., 2011).

<table>
<thead>
<tr>
<th>Species</th>
<th>Seedling (cm)</th>
<th>Sapling (cm)</th>
<th>Pre-reproductive adult (cm)</th>
<th>Reproductive adult (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coryphantha elephantidens Lem.</td>
<td>0-4</td>
<td>4-8</td>
<td>8-12</td>
<td>&gt;12</td>
</tr>
<tr>
<td>Opuntia atropes Rose</td>
<td>1-20</td>
<td>20-80</td>
<td>80-150</td>
<td>&gt;150</td>
</tr>
<tr>
<td>O. pumila Rose</td>
<td>0-200 cm²</td>
<td>200-900 cm²</td>
<td>900-2,500 cm²</td>
<td>&gt;2,500 cm²</td>
</tr>
<tr>
<td>O. velutina</td>
<td>0-8</td>
<td>8-16</td>
<td>16-24</td>
<td>&gt;24</td>
</tr>
<tr>
<td>Pachycereus grandis Rose</td>
<td>1-20</td>
<td>20-80</td>
<td>80-150</td>
<td>&gt;150</td>
</tr>
<tr>
<td>P. weberi Backeb.</td>
<td>1-20</td>
<td>20-150</td>
<td>150-450</td>
<td>&gt;450</td>
</tr>
<tr>
<td>Stenocereus beneckei (Ehrenb.) Buxb.</td>
<td>1-20</td>
<td>20-80</td>
<td>80-150</td>
<td>&gt;150</td>
</tr>
<tr>
<td>S. stellatus (Pfeiffer) Riccob.</td>
<td>1-20</td>
<td>20-80</td>
<td>80-150</td>
<td>&gt;150</td>
</tr>
</tbody>
</table>

Table 1. Life-stage classes of cacti found in six conserved and six perturbed forest plots in the tropical dry forest of Sierra de Huautla, Morelos, Mexico. Note: all size categories were based on the individual height except for Coryphantha elephantidens (diameter) and Opuntia pumila (vegetal cover).
Distance between localities is 8.67 km. Distance among plots ranged from 0.08 to 1.59 km. All cacti found within the plots were counted and measured in January 2007. To analyze population size structure, each individual was assigned to a life-stage class (seedling, juvenile, pre-reproductive, and reproductive adult) based in size and presence of reproductive structures (Martínez-Alvarado, 1985; Table 1).

**Statistical Analysis.** To evaluate cacti density between forest types overall species, a nested Analysis of Variance (ANOVA) was used, where forest type (secondary and conserved) was nested in locality (El Limón and Quila-mula); forest type had three replicates in each locality. Data were transformed with the logarithm base 10 to fulfill ANOVA assumptions (Shapiro Wilk W = 0.95, P < 0.64). To describe the community structure by forest type, dominance/diversity curves were constructed (Wilson, 1991). The Shannon-Wiener diversity index (H’) was calculated using the following formula:

\[ H’ = -\sum_{i=1}^{S} (p_i)\log_2 p_i \]  

(Magurran, 2004)

where H’ is the species diversity index, S the number of species, and p_i is the proportion of individuals from the total number of species that belongs to the species i. To estimate evenness, the Pielou index was used, but first we calculated the maximum evenness with the following formula:

\[ H_{\text{max}} = \log_2 S \]  

(Magurran, 2004)

where H_{\text{max}} is the maximum species richness and S is the species richness observed. When the maximum species richness was calculated, the following formula was used to estimate the Pielou evenness index (Krebs, 1985):

\[ J = H / H_{\text{max}} \]  

(Magurran, 2004)

where J is evenness, H is the observed species richness and H_{\text{max}} is the maximum species richness. H’ was used because it is clearly linked to the dominance/diversity curves (Wilson, 1991). H’ and J were transformed with natural logarithms prior to analysis (Shapiro Wilk W = 0.94, P < 0.51 and W = 0.93, P < 0.35 respectively). H’ values between forest types were compared using t-test for each locality (Zar, 1996).

The Simpson diversity index expresses the probability that two individuals taken randomly from a sample belong to the same species and is calculated as:

\[ D = \sum p_i^2 \]  

(Magurran, 2004)

where p_i is the proportion of the individual of the i species. As D increases, diversity decreases, so it is more commonly used as its inverse (D_{\text{inv}} = 1/D). It takes values from 1 (when there is only one species) to the total species richness of the community (Magurran, 2004).

The Berger-Parker index is a dominance measure that expresses the proportional abundance of the most abundant species:

\[ B = N_{\text{max}} / N \]  

(Magurran, 2004)

where N_{\text{max}} is the number of individuals of the most abundant species and N is the total number of individuals in the community. An increase of B indicates a decrease in evenness and diversity, so its reciprocal is used (B_{\text{rec}} = 1-B); therefore, an increase of B_{\text{rec}} results in an increase in diversity and evenness (Magurran, 2004).

Community size structure (proportion of individuals overall species in each life-stage class) between forest types was compared with Kolmogorov-Smirnov tests and also individually for those species with enough sample size to run statistical analyses. Back-transformed means and standard deviations are shown throughout results. The diversity indexes and species rank curves were calculated using Excel v12.0 and the statistical analyses were carried out using the program R version 2.14.1 (R Development Core Team, 2011).

**Table 2.** Cacti species in conserved and secondary tropical dry forest of Sierra de Huautla, Mexico. Species, life-form, range of height and density are shown. Species were arranged by their density in the conserved forest.

<table>
<thead>
<tr>
<th>Species</th>
<th>Life-form*</th>
<th>Height* (cm)</th>
<th>Conserved</th>
<th>Secondary</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Pachycereus grandis</em></td>
<td>arboreal</td>
<td>500-1,100</td>
<td>146 ± 195.42</td>
<td>134 ± 96.06</td>
</tr>
<tr>
<td><em>Opuntia pumila</em></td>
<td>shrub-like</td>
<td>50-60</td>
<td>23.33 ± 49.54</td>
<td>18 ± 38.26</td>
</tr>
<tr>
<td><em>Opuntia puberula</em></td>
<td>shrub-like prostrate</td>
<td>50-60</td>
<td>13.33 ± 15.93</td>
<td>95.33 ± 95.01</td>
</tr>
<tr>
<td><em>Coryphantha elephantidens</em>†</td>
<td>globose</td>
<td>12-14</td>
<td>2.67 ± 4.13</td>
<td>48.67 ± 98.12</td>
</tr>
<tr>
<td><em>Opuntia velutina</em></td>
<td>arboreal</td>
<td>200-350</td>
<td>2.67 ± 4.84</td>
<td>104.67 ± 83.65</td>
</tr>
<tr>
<td><em>Stenocereus stellatus</em>†</td>
<td>arboreal</td>
<td>100-350</td>
<td>2 ± 3.35</td>
<td>13.33 ± 9.35</td>
</tr>
<tr>
<td><em>Stenocereus beneckei</em>†</td>
<td>shrub-like</td>
<td>100-200</td>
<td>0.67 ± 1.63</td>
<td>7.33 ± 8.16</td>
</tr>
<tr>
<td><em>Opuntia atropes</em></td>
<td>arboreal</td>
<td>100-300</td>
<td>-</td>
<td>2.67 ± 4.13</td>
</tr>
<tr>
<td><em>Pachycereus weberi</em>†</td>
<td>arboreal</td>
<td>700-1,100</td>
<td>-</td>
<td>1.33 ± 3.27</td>
</tr>
</tbody>
</table>

* Based on Martínez (1985)
† mostly endemic (Hernández and Godínez, 1994)
Results

In total, 943 individuals from nine species were registered, measured and identified (Table 2). Two large arboreal (~11 m height), three short arboreal (~3 m height), one large shrub (~2 m height), and three small size life-forms (shrubs, prostrate and globose < 60 cm height) were recorded. All species were registered in both forest types except for Opuntia atropes and Pachycereus weberi, which were found exclusively in the secondary forest (Table 2). The large arboreal species Pachycereus grandis was the most abundant in conserved (70.65% of individuals) and secondary forest (31.75%; Table 2, 3). After P. grandis, the short arboreal O. velutina (27%) and the prostrate O. puberula (22.59%) were the species with the highest abundance in the secondary forest, whereas the small shrub O. pumila (19%) was the most abundant in the conserved forest (Table 2, 3). In average and taking into account all species and life-stage classes, there was a higher cacti density in the secondary forest (309.14 ± 2.62 ind./ha) than in the conserved forest (119.69 ± 3.85 ind./ha). However, density was statistically similar in both forest types (F(2,8) = 1.94, P > 0.10).

The dominance/diversity curve showed that there was higher richness in the secondary than in the conserved forest. The dominance/diversity curve was steeper in the conserved forest (Figure 1, Table 4) suggesting higher richness and evenness in the secondary forests of both localities. The Shannon-Wiener, inverse Simpson and the reciprocal of Berger-Parker diversity index confirmed the result of the curves: a significantly higher indexes were found in the secondary forest (H’ > 1.39, Dinv > 2.88, Brec > 0.53) than in the conserved forest (H’ > 0.42, Dinv > 1.23 Brec > 0.82) from two localities (t(2,8) = 4.8, P < 0.005) due to both, a higher richness and a higher evenness in the secondary forest (Figure 1, Table 4).

Size structure. At community level, the proportion of individuals in smaller life-stage classes (seedlings and juveniles) was two times higher in the secondary forest compared to the conserved forest, whereas the proportion of reproductive adults was two times higher in the conserved forest (Table 5). At community level, size structure was significantly different by forest type (Dmax = 310, P < 0.001; Table 5).

Two species, the large arboreal Pachycereus grandis and the small shrub Opuntia pumila had enough individuals at all life-stage classes and in both forest types to run statistical analyses at species level. The population size structure of P. grandis showed that the proportion of individuals in the seedlings and juveniles life-stage classes were 18 and three times higher, respectively in the secondary forest than in the conserved forest; whereas the proportion of individuals in pre-reproductive and reproductive adults classes were al-

Table 3. Abundance (N) and average size (± standard deviation [SD]) of cactus species in twelve 50 × 50 m plots in conserved and secondary tropical dry forest at two localities (El Limón and Quilamula) at Sierra de Huautla, Morelos, Central Mexico. * Height (cm) for all species except when indicated (§).

<table>
<thead>
<tr>
<th>Species</th>
<th>El Limón Plot</th>
<th>Quilamula Plot</th>
<th>N</th>
<th>Size ± SD *</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Conserved</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coryphantha elephantidens</td>
<td>2</td>
<td>1</td>
<td>4</td>
<td>11.1 ± 2.7</td>
</tr>
<tr>
<td>Opuntia atropes</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>Opuntia puberula</td>
<td>6</td>
<td>1</td>
<td>20</td>
<td>123.0 ± 41.7</td>
</tr>
<tr>
<td>Opuntia pumila</td>
<td>0</td>
<td>0</td>
<td>59</td>
<td>15.1 ± 9.6</td>
</tr>
<tr>
<td>Opuntia velutina</td>
<td>0</td>
<td>1</td>
<td>4</td>
<td>156.5 ± 191.1</td>
</tr>
<tr>
<td>Pachycereus grandis</td>
<td>88</td>
<td>109</td>
<td>219</td>
<td>467.8 ± 43.0</td>
</tr>
<tr>
<td>Pachycereus weberi</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>Stenocereus beneckei</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>130</td>
</tr>
<tr>
<td>Stenocereus stellatus</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>175.0 ± 114.6</td>
</tr>
<tr>
<td><strong>Secondary</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coryphantha elephantidens</td>
<td>63</td>
<td>5</td>
<td>74</td>
<td>10.1 ± 3.1</td>
</tr>
<tr>
<td>Opuntia atropes</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>320.0 ± 43.2</td>
</tr>
<tr>
<td>Opuntia puberula</td>
<td>47</td>
<td>58</td>
<td>143</td>
<td>2,619.1 ± 3,811.6 §</td>
</tr>
<tr>
<td>Opuntia pumila</td>
<td>1</td>
<td>1</td>
<td>27</td>
<td>17.0 ± 7.8</td>
</tr>
<tr>
<td>Opuntia velutina</td>
<td>53</td>
<td>47</td>
<td>167</td>
<td>86.6 ± 77.2</td>
</tr>
<tr>
<td>Pachycereus grandis</td>
<td>21</td>
<td>41</td>
<td>201</td>
<td>148.2 ± 80.9</td>
</tr>
<tr>
<td>Pachycereus weberi</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>670.0 ± 608.1</td>
</tr>
<tr>
<td>Stenocereus beneckei</td>
<td>3</td>
<td>4</td>
<td>11</td>
<td>89.6 ± 57.8</td>
</tr>
<tr>
<td>Stenocereus stellatus</td>
<td>5</td>
<td>0</td>
<td>5</td>
<td>214.6 ± 186.1</td>
</tr>
</tbody>
</table>
most three times lower in the secondary forest compared to the conserved forest (n = 66 in the secondary forest and n = 219 in the conserved forest, Figure 2A). The population size structure of *P. grandis* showed significant differences between forest types ($D_{\max} = 114, P < 0.001$).

The population size structure of *Opuntia pumila* showed that the proportion of seedlings and adults was higher in the conserved forest compared to the secondary forest, whereas the proportion of juveniles and pre-reproductive adults were higher in the secondary forest (Figure 2B; Table 5). However, the population size structure of *O. pumila* was statistically similar between forest types ($D_{\max} = 6, P > 0.1$).

**Discussion**

Richness, diversity and evenness of the community of cacti in the dry forest of Sierra de Huautla increased in sites under chronic disturbance whereas density was not significantly affected. Size structure at community level in the conserved forest showed an increase of individuals with size, whereas in the secondary forest size structure showed an uneven distribution of individuals, as in populations that regenerate by pulses. Further, in the perturbed forest, there was a lower proportion of adults and they were, in average, smaller than those recorded in the conserved forest.

**Community structure.** Studies at the population level have shown positive, negative or neutral effects of disturbance in cacti density. Some species (called ruderal cacti *sensu* Martorell and Peters, 2005) may benefit from moderate levels of human activities as the globose species *Coryphantha werdermannii* in Coahuila, North Mexico (Portilla-Alonso and Martorell, 2011) and the cylindrical cactus *Echinocereus lindsayi* in Baja California, Mexico (Martorell et al., 2012). In the present study, further analysis at species level revealed that *Opuntia velutina* and *O. puberula* were significantly more abundant in the secondary forest under chronic disturbance compared to the conserved forest ($t_{(1,10)} = 2.6, P < 0.05$ and $t_{(1,10)} = 6.4, P < 0.0001$, respectively). This is in agreement with a study in the dry forest of Yucatan, Mexico, where it was found that the globose cactus *Mammillaria gaumeri* benefits from an increase of light after moderate disturbance of the forest canopy (Cervera et al., 2007). Further, four species evaluated in the present study showed similar abundance in secondary and conserved forest: *Coryphantha elephantidens* ($t_{(1,10)} = 1.8, P > 0.1$), *Pachycereus grandis* ($t_{(1,10)} = 0.9, P > 0.4$), *O. pumila* ($t_{(1,10)} = 0.6, P > 0.5$) and *Stenocereus beneckei* ($t_{(1,10)} = 1.8, P > 0.1$). This was in agreement to the neutral effect of disturbance found for *Mammillaria crucigera* and *M. huizilopochtli* in the Tehuacan Valley, Puebla, Mexico (Martorell and Peters, 2009). More than 60 % of the cactus species occurring in this dry forest showed neutral or positive effect of disturbance; this seems to be reflected in the neutral effect of disturbance in the density of the entire cacti community considering all life-stage classes.

Cacti diversity, richness, and evenness increased in the secondary forest under chronic disturbance. These results are in agreement with what is known for smaller life-forms, as grasses and herbs experiencing faster succession after disturbance. For example, in mountain grasslands of Argentina (Nai-Bregaglio et al., 2002) and in the tropical dry forest of...
Figure 2. Population structure of A) *Pachycereus grandis* and B) *Opuntia pumila* in conserved (black bars) and secondary (white bars) tropical dry forest of Sierra de Huautla, Mexico. Life-stage classes are S = seedlings, J = juveniles, PRA = pre-reproductive adults and RA = reproductive adults.

Sierra de Huautla, Mexico (De la O-Toris et al., 2012), a higher diversity and richness of herbs and grasses was found in disturbed areas compared to conserved ones (but see Floyd et al., 2003). In this study, seven out of nine cactus species and more than 55% of individuals registered in both forest types were ≤ 3 m in height. Therefore, the higher richness, diversity and evenness of cacti in the secondary forest may reveal the response to disturbance of the smaller life-forms, similar to what happen to herbs and grasses in natural grasslands under chronic disturbance by livestock.

It is expected that cactus species inhabiting the secondary forest be a subset of those occurring at the conserved old forest. However, in this study, *Opuntia atropes* and *Pachycereus weberi* were only found in the secondary forest from one of the localities evaluated (Quilamula). These two species are used as food and also for feeding cattle (Luna-Morales and Aguirre, 2001; López-Acosta et al., 2008; Arias and Terrazas, 2009). *Opuntia atropes* (commonly named White nopal) is reported as native in various states in central Mexico (Guanajuato, Guerrero, Jalisco, Mexico, Michoacan and Morelos; Instituto de Biología, 2013) and it is also used in traditional medicine and to build living fences (López-Acosta et al., 2008). Further, *O. atropes* is one of the two species more frequently used as vegetable and therefore it has been cultivated probably since prehistoric times (Callen, 1967). *P. weberi* is also used as food for

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<th>Table 5. Proportion of cactus individuals by life-stage classes in secondary and conserved tropical dry forest of Sierra de Huautla, Mexico. Life-stage classes (Seedlings, Juveniles, Pre-reproductive adults [Pre-R] and Adults) were based in size (Table 1).</th>
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<tbody>
<tr>
<td>Species</td>
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<td>Seedlings</td>
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<tr>
<td>All species</td>
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<tr>
<td><em>Coryphantha elephantidens</em></td>
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<td><em>Opuntia atropes</em></td>
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<td><em>Pachycereus grandis</em></td>
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cattle and their seeds used to elaborate flatbread (tortillas), sauces (mole), and also for construction (Luna-Morales and Aguirre, 2001; Arias and Terrazas, 2009). The presence of *P. weberi* has been registered in ancient homegardens and is usually tolerated in agricultural lands (Luna-Morales and Aguirre, 2001). Many plant species widely used by people in the dry forest grow in sites under chronic disturbance (De la O-Toris *et al*., 2012; Maldonado *et al*., 2013). The multiples uses of these two species and their presence in low density only in the secondary forest suggest that they could be heavily extracted at both forest and their populations have not recovered. In this study, the presence of two more cactus species in the secondary forest and a similar density of individuals between forest types increased community evenness; this forest structure seems to be the result of some cactus species been favoured by microenvironmental conditions whereas others were heavily extracted.

**Size structure.** In the conserved forest, the proportion of individuals increased with size in all the species, which is a structure seldom registered for cactus species (reviewed in Godínez-Álvarez *et al*., 2003). In the arid region of Cuatro Cienegas, Coahuila, North Mexico, an increment of individuals with size was registered for the globose species *Coryphantha werdermannii*, at sites with very low disturbance (Portilla-Alonso and Martorell, 2011). A higher proportion of individuals in the larger than the smaller size-classes may be related to recruitment limitation (Nathan and Muller-Landau, 2000); recruitment limitation may occur at saturated undisturbed communities with a continuous canopy due to unsuitable microenvironmental conditions for seed germination and recruitment (see e.g. Clark *et al*., 1998). On the other hand, changes in size structure of cactus species due to disturbance has been registered in forest ecosystems. In the dry forest of the state of Veracruz, southeast Mexico, the cylindrical cactus *Mammillaria eriacantha* showed a decrease of individuals with size in most of the fragments evaluated (Valencia-Díaz *et al*., 2012); that size structure is related to continuous regeneration (Godínez-Álvarez *et al*., 2003). However, at the most disturbed fragment, the population of *M. eriacantha* showed an uneven size structure (Valencia-Díaz *et al*., 2012). In this study, the large arboreal cactus *Pachycereus grandis* showed a change in size structure with disturbance similar to that registered for the entire cacti community, whereas the shrub-like cactus *Opuntia pumila* showed a similar size structure in both forest types: the size structure related to constant regeneration. Therefore, a change in the size structure in the entire community and in the population of *P. grandis* due to disturbance may be the result of better recruitment conditions in the secondary forest.

In the conserved forest, the proportion of adults overall species was two times higher than the one recorded in the secondary forest, according to our hypothesis. These results are in line with others studies at population level; for example for *Ferrocactus histrix* (del Castillo, 1987) and *Coryphantha werdermannii* (Portilla-Alonso and Martorell, 2011), a higher proportion of adults were found at the sites with less chronic disturbance. The proportion of adults under conditions of chronic disturbance may decrease due to high mortality of small size-classes and slow growth rates (Jiménez-Sieerra *et al*., 2007; Alba-García, 2011; Martorell *et al*., 2012). Also, some studies have showed that even when recruitment is higher in grasslands, cacti growth faster in sites denuded of grasses (Mandujano *et al*., 2001) because belowground competition (Briones *et al*., 1996). This seems to be supported by the average height of adults of *Pachycereus grandis* in this study which was three times higher in the conserved forest (468 ± 43) compared to the secondary one (148 ± 81; Table 3). In the conserved forest, the higher proportion of larger adults was probably due to higher growth rates and lower recruitment. Few studies have explored the regeneration mechanisms of dry forest trees (reviewed in Quesada *et al*., 2009), whereas cactus species, with a variety of life-forms, are seldom included in the studies of woody species or totally excluded from the ones evaluating understory vegetation. Therefore, the mechanisms that explain the size structure of cactus species in the dry forest and its changes with disturbance needs to be further explored.

**Implications for conservation and restoration of biodiversity in dry forests.** Secondary forests that resulted from land abandonment after anthropogenic activities now cover more areas than conserved forest (Chazdon *et al*., 2007). These forests are important reservoir of biodiversity and because cactus populations may have different responses to disturbance, additional studies should be carry out to support management projects for their conservation (Martorell and Peters, 2009; Ureta and Martorell, 2009; González *et al*., 2013). In this study, 18 cactus species previously reported for Sierra de Huautla Biosphere Reserve (Martínez-Alvarado, 1985; CO-NANP, 2005) were not recorded in our plots probably because some of them are specialists of habitats not covered (e.g. rupeicolous species). In these dry forests, some cactus species are specialists of uncommon microhabitats; conservation in those areas where many microhabitats are represented is crucial for cacti conservation (see e.g. Gibson *et al*., 2011).

Studies of succession have shown that vegetation that tolerates very dry conditions may successfully invade degraded landscapes (Ewel, 1980). We believe this may be the case for *Opuntia puberula* and *O. velutina* which have densities of one order of magnitude higher in the secondary forest compared to the conserved forest. However, the higher densities in the secondary forest may also be the result of asexual reproduction. For example, in the Southern Chihuahuan Desert, sexual reproduction of *O. rastrera* is lower than asexual reproduction, because the latter is enough to maintain these populations (Mandujano *et al*., 1996, 2001). Therefore, further studies should be conducted in the dry
forest to evaluate if densities of *Opuntia* spp. are higher due to microclimatic conditions of secondary forests or to their asexual reproductive potential.

**Conclusions**

Chronic disturbance in this secondary forest during more than 30 years changed size structure of cactus species resulting in less adult individuals of reduced size. Under chronic disturbance, animals that depend on large arboreal cacti for food (i.e., bats, ants, bees, and butterflies) and perches or refuge (i.e., woodpecker) may move out affecting negatively many plant-animal interactions and the maintenance of diversity derived from them (Noy-Meir, 1974). For example, disturbed dry forests are usually dominated by trees with dry fruits dispersed by wind (Martínez-Garza et al., 2011) and therefore, these areas are not attractive to frugivores. One option to favour the coexistence of economic activities and biodiversity is to exclude fragments or medium size dry forests are usually dominated by trees with dry fruits dispersed by wind (Martínez-Garza et al., 2011) and therefore, these areas are not attractive to frugivores. One option to favour the coexistence of economic activities and biodiversity is to exclude fragments or medium size

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