Research Article

Natural hybridization and genetic and morphological variation between two epiphytic bromeliads

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Abstract. Reproductive isolation is of fundamental importance for maintaining species boundaries in sympatry. Here, we examine the genetic and morphological differences between two closely related bromeliad species: Vriesea simplex and Vriesea scalaris. Furthermore, we examined the occurrence of natural hybridization and discuss the action of reproductive isolation barriers. Nuclear genomic admixture suggests hybridization in sympatric populations, although interspecific gene flow is low among species in all sympatric zones (Nm < 0.5). Thus, morphological and genetic divergence (10.99 %) between species can be maintained despite ongoing natural hybridization. Cross-evaluation of our genetic and morphological data suggests that species integrity is maintained by the simultaneous action of multiple barriers, such as divergent reproductive systems among species, differences in floral traits and low hybrid seed viability.

Keywords: Atlantic Forest; Bromeliaceae; floral traits; hybrids; reproductive barriers; species integrity; sympatric.

Introduction

Natural hybridization is an important process in plant evolution. It has been estimated that 30–70 % of all flowering plant species have hybridization events in their phylogenetic histories (Ehrlich and Wilson 1991; Rieseberg 1995; Soltis and Soltis 2009). Therefore, hybrid zones are interesting models for studying the evolution of reproductive barriers, the role of selection in maintaining species differences and how phenotypic traits differ between hybridizing populations (Abbott et al. 2013; Arnold 2014).

The degree of the reproductive isolation barrier among related species is an important factor that influences the genetic integrity of a species and the probability of forming a hybrid (Coyne and Orr 2004). One of the key points in evolutionary biology is to determine how reproductive barriers limit introgressive gene flow and hybridization (Coyne and Orr 2004). Congruent hybridization patterns can either unify populations of species or preserve two populations of species with different allele frequencies (Servedio and Kirkpatrick 1997). Asymmetric gene flow is important to understanding the factors that determine the maintenance of species identity. Some of these
underlying factors include variations in the reproductive system (Hu 2015) and flower size (floral traits). Flower size may also be a major structural barrier to hybridization that is generally asymmetrical (Williams and Rouse 1988). This feature is particularly important because flower sizes generally differ between hybridizing species (Field et al. 2011).

Reproductive isolation barriers can function prior to mating (premating), after mating but before zygote formation (postmating–prezygotic) and after zygote formation (postzygotic) (Ramsey et al. 2001, 2002; Moccia et al. 2007; Pinheiro et al. 2010; Palma-Silva et al. 2011; Brys et al. 2013; Briscoe Runquist et al. 2014; Marques et al. 2014; Pinheiro et al. 2015; Twyford et al. 2015).

Mating systems are recognized as key barriers to reproductive isolation (Jain 1976; Coyne and Orr 2004), although their role in speciation and species cohesion is not completely understood, especially for plants with mixed mating systems (Hu 2015). The floral adaptations that allow autonomous selfing are assumed to offer effective mechanical protection against heterospecific mating and thus to contribute to reproductive isolation (Wright et al. 2013). Empirical support for selfing as a reproductive barrier was reported for different plant groups (Fishman and Wyatt 1999; Wendt et al. 2002; Fishman and Stratton 2004; Lowe and Abbott 2004; Martin and Willis 2007; Matallana et al. 2010). In addition, sympatric species with a shared generalized floral morphology and pollinator community, intense competition for pollination and fitness costs related to hybridization may select for floral traits that contribute to prezygotic isolation (Servedio and Noor 2003). For example, Beans (2014) suggests that the divergence of floral traits in sympatric and allopatric populations may evolve in response to competition for pollinator resources or in response to costs associated with sharing pollinators with other species.

Neotropical plant radiations provide a perfect system for examining potential gene flow among closely related species. The Bromeliaceae, in particular, have been used as a model group in studies of evolution of reproductive isolation and speciation in Neotropical regions (Schulte et al. 2010; Palma-Silva et al. 2011, 2015, 2016a, b; Wagner et al. 2015; Lexer et al. 2016; Zanella et al. 2016), mainly due to its high diversity and recent adaptive radiation (Benzing 2000; Givnish et al. 2011, 2014).

Here we investigated natural hybridization and the factors involved in the maintenance of phenotypic and genetic differentiation between two recently diverged species (Barfuss et al. 2016) that occur in the Brazilian Atlantic Forest: Vriesea simplex and Vriesea scalaris. The two species are self-compatible (Matallana et al. 2010), but their reproductive systems are divergent, ranging from predominantly outcrossing and pollinator dependent in V. simplex to highly selfing in V. scalaris (Neri et al. 2017). Both species exhibit similar pollination syndromes, their flowers are visited by hummingbirds (Phaethornis eurynome and Ramphodon naevis) and overlap in flowering time occurs in sympathy (Wendt et al. 2008). In addition, artificial hybrids (F1) were obtained via manual crossing experiments involving these two taxa (Neri et al. 2017), confirming interspecific compatibility. Despite reports of artificial cross-compatibilities and observation of putative hybrids in the field, the extent of gene exchange between these two species in natural populations has not been investigated.

We investigated four allopatric and three sympatric populations using a combined set of microsatellite markers and multivariate analyses of morphology to answer the following questions: (i) How do genetic and morphological differences between sympatric and allopatric populations contribute to reproductive isolation between V. simplex and V. scalaris? (ii) Do V. simplex and V. scalaris hybridize in the wild, as suggested by artificial crosses and observations of putative hybrids in the field? If yes, are the patterns of hybridization and gene flow (migration) similar or asymmetrical across sympatric populations? (iii) What is the importance of the action of different prezygotic and postzygotic barriers in maintaining species integrity?

Methods

Population samples and DNA extraction

Vriesea simplex and V. scalaris are epiphytic species that occur in mesophlic environments and well-preserved habitats with high humidity in the Brazilian Atlantic rainforest (Fig. 1). Vriesea simplex has a narrow distribution (Bahia, Espírito Santo, Rio de Janeiro and São Paulo), whereas V. scalaris has a widespread distribution (Pernambuco to Rio Grande do Sul states; Forzza et al. 2015). We sampled three hybrid zones: Santa Lucia (EBS; 108 individuals), Ruschi (RUS; 40 individuals) and Duas Bocas (RDB; 33 individuals); two allopatric collection locales of V. simplex: Guapimirim (GUA; 16 individuals) and Soberbo (SOB; 18 individuals); and two allopatric collection locales of V. scalaris: Peri (PER; 20 individuals) and Sincorá (SIN; 10 individuals). In total, 255 flowering or fruiting plants
were sampled (Table 1; Fig. 1). Voucher information for each collection locale and species is given in Supporting Information—Table S1. The identified hybrids and their intermediate morphology are described in Supporting Information—Table S2. Fresh leaves from each individual were collected and stored in silica gel. Total genomic DNA was extracted using a Invisorb Spin Plant Mini Kit (Stratec Biomedical AG, Birkenfeld, Germany) according to the manufacturer’s instructions.

**Nuclear microsatellite markers and genotyping**

To study the patterns of genetic diversity and genomic admixture in sympatric and allopatric populations, we used 15 nuclear microsatellite loci (SSR) previously developed for other bromeliad species. Six loci were isolated from *V. simplex* (Vs1, Vs2, Vs8, Vs9, Vs10 and Vs19; Neri et al. 2015), six loci from *V. gigantea* (VgB01, VgB10, VgB12, VgG02, VgG04 and VgG05; Palma-Silva et al. 2007), two loci from *Tillandsia fasciculata* (E6 and E6b; Boneh et al. 2003) and one locus from *Aechmea caudata* (Ac01; Goetze et al. 2013). For each SSR, the forward primers were synthesized with an M13 tail (5′-CACGACGTTGTAAAACGAC-3′) to allow for marking and multiplexing fluorescent dyes during the amplification and genotyping procedures. All PCR amplification reactions were performed in a Thermal Cycler (Applied Biosystems, Foster City, CA, USA) following the protocol described by Palma-Silva et al. (2007). The microsatellite alleles were resolved on a 3500 DNA Analyzer automated sequencer (Applied Biosystems) and sized against the GeneScan 500 LIZ molecular size standard (Applied Biosystems) using GENEMARKER Demo version 1.97 software (SoftGenetics, State College, PA, USA). Microchecker software (Van Oosterhout et al. 2004) was used to check for null alleles.

Figure 1. Map of localities of samples collected from sympatric and allopatric populations of *Vriesea simplex* and *V. scalaris* in the Atlantic rainforest used for study of hybridization and morphometric analysis. *Vriesea simplex*, ‘sim’; *V. scalaris*, ‘sca’. For abbreviations of populations, see Table 1.
Statistical analysis

**Patterns of genomic admixture for hybrid detection.** Bayesian analysis was performed in STRUCTURE 2.3.2 software (Pritchard et al. 2000) and carried out under the admixture model assuming independent allele frequencies, using a burn-in period of 100,000, run length of 500,000 and 10 replicates per K ranging from 1 to 10 with all populations in the data set. Using the method proposed by Evanno et al. (2005), which is based on an ad hoc measure of ΔK, we determined the highest number of clusters (K) as K = 2, corresponding to the two species. We used the K = 2 model and 10 replicates per K because we assumed that the two species contributed to the gene pool of the sample. Allopatric populations of each species were used as reference samples of pure V. simplex and V. scalaris. Follow-up analyses were performed separately for each hybrid zone, in each case including the specimens from the allopatric populations as reference samples for each species. We investigated the genetic structural patterns of each hybrid zone separately because the allelic frequencies within the populations of each species were different (see Table 3), particularly because of the high and variable selfing rates observed for V. scalaris (Table 3). It is also important to analyse hybrid zones separately when gene flow is restricted among populations, leading to high divergence and different genomic architecture among hybrid zones (Pinheiro et al. 2010; Marques et al. 2014; Twyford et al. 2015; Zanella et al. 2016). Following the procedure described by Burgarella et al. (2009), STRUCTURE was used to classify individuals among the two parental species and hybrids, using a threshold value of q = 0.75; individuals with q < 0.75 remained unassigned.

**Genetic diversity.** Populations and loci were characterized for V. simplex, V. scalaris and their hybrids based on number of alleles, allelic richness, variance in
allele size, observed and expected heterozygosity and inbreeding coefficient \((F_{IS})\) using the MSA (Dieringer and Schlötterer 2003) and FSTAT softwares (Goudet 1995). We also estimated the private alleles of each species and hybrids using GenAlEx software (Peakall and Smouse 2006).

Departures from Hardy–Weinberg equilibrium (HWE) were tested using the web-based GENEPOP 3.5 software (Raymond and Rousset 1995). Using the \(F_{IS}\) we calculated the apparent outcrossing rate \((t_p)\), according to the following formula: \(t_p = (1 - F_{IS})/(1 + F_{IS})\) (Goodwillie et al. 2005). We assume that predominantly selfing populations have \(t_p \leq 0.2\), mixed systems have \(0.2 < t_p \leq 0.8\) and predominantly outcrossing populations have \(t_p > 0.8\) (Goodwillie et al. 2005).

Partitioning of genetic diversity within and among \(V.\) simplex and \(V.\) scalaris groups was evaluated by an analysis of molecular variance (AMOVA) implemented in the software ARLEQUIN 3.1 (Excoffier et al. 2005). Principle coordinate analysis (PCoA) was used on the entire data set to visualize genetic differences between species and allopatric and sympatric populations, and to examine the genetic status of plants along the contact zone, implemented in the GenAlEx program (Peakall and Smouse 2006).

**Effective population size and migration rate.** We calculated the effective population size \((N_e)\) of \(V.\) simplex and \(V.\) scalaris because we expected that introgression might occur more intensely in populations with a smaller effective population size. Effective migration rates \((N_{m})\) were estimated between pairs of sympatric populations of \(V.\) simplex and \(V.\) scalaris following a coalescent theory and maximum-likelihood-based approach using MIGRATE 3.0.3 software (Beerli and Felsenstein 1999). The computations were carried out on the infinite allele model (Kimura and Crow 1964). Effective population size values were estimated from theta values by assuming a microsatellite \(\mu\) rate of \(10^{-3}\) per gamete per generation (Zhang and Hewitt 2003).

**Sampling and morphometric analysis.** Morphometric analyses of vegetative and reproductive traits of pure species (individuals of both species) and hybrids were performed in two sympatric populations, three allopatric collection locales of \(V.\) simplex and five allopatric collection locales of \(V.\) scalaris (Table 1; Fig. 1). All individuals included in morphometric analyses were identified as parental species or hybrids using SSR genotypes based on STRUCTURE analysis, as described above.

Individuals were collected and measured based on availability and accessibility. In total, we measured 139 flowering specimens from both parental species and hybrids (Table 1). Twenty-four quantitative species and hybrids (Table 1). Twenty-four quantitative traits (6 vegetative and 18 reproductive; Table 5) were measured using callipers. The flowers were collected and preserved in 70% ethanol. A discriminant analysis (canonical variance analysis; CVA) was performed in the STATISTICA8 program for Windows 4.2 (StatSoft 1993) to test the partition among predefined clusters (\(V.\) simplex allopatric, \(V.\) scalaris allopatric, \(V.\) simplex sympatric, \(V.\) scalaris sympatric and hybrids) and to identify traits that contribute most to species discrimination.

We used the most significant traits for species discrimination in CVA to examine the extent of variation among sympatric and allopatric collection locales between species. These comparisons were performed using analysis of variance (ANOVA), followed by Tukey's test through the general linear model. These statistical analyses were performed using R software (Core Team 2015).

**Results**

**Genetic composition of hybrid zones**

Genomic admixture analysis with Bayesian STRUCTURE results for sympatric populations indicated hybridization between \(V.\) simplex and \(V.\) scalaris, with a total of 22 hybrids identified among 252 individuals sampled (12% of the total individuals sampled in sympatric populations; threshold: \(0.10 < q < 0.90\), Fig. 2). The EBS population had 12 hybrids among 108 individuals, the RUS population had 8 hybrids among 40 individuals and the RDB population had only 2 hybrids among 33 individuals. Most hybrids identified in STRUCTURE were not assigned into any hybrid class using NEWHYBRIDS. In total, NEWHYBRIDS was able to classify 10 hybrid individuals, all as \(F_{2}\), in the EBS population, and 2 in the RUS population. In the RDB population, no hybrid was classified by NEWHYBRIDS (Fig. 3).

**Nuclear microsatellite diversity**

Levels of genetic diversity differed strongly between species (Table 2), with \(V.\) simplex having a total of 262 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) scalaris having a total of 92 alleles (ranging from 3 to 18 alleles per locus). Hybrids presented a total of 110 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) simplex having a total of 262 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) scalaris having a total of 92 alleles (ranging from 3 to 18 alleles per locus). Hybrids presented a total of 110 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) simplex having a total of 262 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) scalaris having a total of 92 alleles (ranging from 3 to 18 alleles per locus). Hybrids presented a total of 110 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) simplex having a total of 262 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) scalaris having a total of 92 alleles (ranging from 3 to 18 alleles per locus). Hybrids presented a total of 110 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) simplex having a total of 262 alleles (ranging from 6 to 40 alleles per locus) and \(V.\) scalaris having a total of 92 alleles (ranging from 3 to 18 alleles per locus).
V. scalaris (Table 3). We observed a higher number of private alleles for V. simplex, with 128 private alleles (out of 262 alleles), than in V. scalaris, with 6 private alleles (out of 92 alleles), or in hybrids, with 7 private alleles (out of 110 alleles).

The number of alleles in populations ranged from 102 to 170 in V. simplex and from 29 to 42 in V. scalaris (Table 3). Population-level $F_{IS}$ values departed significantly from HWE in almost all populations. The $F_{IS}$ values were higher with significant heterozygote deficits more prevalent in V. scalaris ($F_{IS} = 0.511$) than in V. simplex ($F_{IS} = 0.179$), consistent with differences in their reproductive systems (Neri et al. 2017). In agreement with mating system variation between species, the apparent outcrossing rates ($t_o$) were higher for V. simplex, ranging from 0.658 to 0.845, than for V. scalaris, ranging from 0.158 to 0.753 (Table 3). Despite this, all $t_o$ values for both species were between 0.2 and 0.8, suggesting mixed systems for both species (Table 3). In addition, V. simplex $t_o$ values were similar among sympatric and allopatric populations, but V. scalaris sympatric populations had lower $t_o$ values than allopatric populations, indicating selfing rates may be higher in all sympatric populations (Table 3). Hybrids showed, on average, an intermediate genetic diversity index compared to purebred species.

### Genetic differentiation and nuclear migration rates between species

AMOVA results showed that genomic differentiation between species was low (10.99 %), but still highly significant ($P < 0.001$; Table 4). The separated AMOVA model for each species indicated lower genetic structure among populations of V. simplex ($F_{ST} = 0.069; P < 0.001$) than for V. scalaris ($F_{ST} = 0.416; P < 0.001$) (Table 4). The PCoA produced two defined groups of pure V. simplex and pure V. scalaris. The hybrids did not form an intermediate group and several hybrids were grouped with V. scalaris (Fig. 4).

The maximum-likelihood-based estimates of the effective numbers of migrants ($N_{m}$) for sympatric populations of V. simplex and V. scalaris were very low among species, suggesting restricted interspecific gene flow. Although low, interspecific migration rates were
asymmetric towards V. simplex, with larger $N_e$ values from V. scalaris into V. simplex (Fig. 5). The $N_e$ sizes were larger for V. simplex (ST, $N_e = 3080$; RUS, $N_e = 5972.50$; RBD, $N_e = 5362.50$; GUA, $N_e = 3.60$; and SOB, $N_e = 4.21$) than for V. scalaris (ST, $N_e = 1041.55$; RUS, $N_e = 395.05$; RBD, $N_e = 404.97$; SIN, $N_e = 426.95$; and PER, $N_e = 115.65$).

### Morphometric analysis of vegetative and reproductive traits among species

CVA morphometric analysis of 24 characters (18 reproductive and 6 vegetative) distinguished two well-defined groups, consistent with parental species identified using microsatellites. Hybrids did not present intermediate trait grouping with one or the other species (Fig. 6). The first CVA axis accumulated 97 % of the total variation among clusters (Fig. 6). The six variables that contributed most to this axis were floral traits: scape length; length and width of the floral bract; length and width of the bract floral scape; and pistil length (Table 5). Additionally, three out of the six floral traits (floral bract length, floral bract width and scape bract width) were significantly higher in sympatric populations than in allopatric populations of V. simplex (Fig. 7). Scape length was significantly lower in sympatric than in allopatric populations of V. simplex (Fig. 7).

### Discussion

In this study, we investigated the potential evolutionary mechanisms associated with maintenance of reproductive species barriers between V. simplex and V. scalaris by examining morphological and genetic variation of these species in the Brazilian Atlantic Forest. Our results revealed three important points: (i) these species can be considered two distinct taxa, supported by genetic and morphological data, even with the occurrence of natural hybridization; (ii) divergent levels of genetic diversity (lower in V. scalaris) and $F_{IS}$ (higher in V. scalaris) are in agreement with reproductive system variation in these species (Neri et al. 2017), with predominance of selfing in V. scalaris and outcrossing in V. simplex; (iii) variation in floral characters among sympatric and allopatric populations occurring only in the outcrosser V. simplex, suggests sympatric floral display in V. simplex tends to be showier than in allopatry. The variations in reproductive systems and floral traits may be potential prezygotic...
barriers. Although incomplete, the combination of prezygotic barriers (divergent mating system and floral display) together with postzygotic barriers (inviable hybrid seeds), may act to maintain the morphologic and genetic integrity of these incipient species, even in the presence of hybridization. The different approaches used in this study provide information on the processes involved in maintaining the integrity of correlated species.

## Genetic and morphological differentiation and species integrity

Genetic differentiation between species (AMOVA–$F_{CT} = 10.99\%$, P-value < 0.001), differences in distributions of allele frequencies between species (PCoA; Fig. 4) and low levels of interspecific gene flow ($N_{me} = 0.05–0.24$; Fig. 5) in sympatric populations of Vriesea suggest these species are indeed independent evolutionary units. Despite the differentiation between species supported by genetic and morphologic data, ancestral polymorphisms or recent gene flow (Coyne and Orr 2004) could still be present between these sister species (Barfuss et al. 2016). Ancestral polymorphism sharing is likely due to recent species divergence in the Vriesea genus (Givnish et al. 2014; Gomes-da-Silva 2013; Barfuss et al. 2016) and/or incomplete lineage sorting (e.g. Costa et al. 2009; Zanella et al. 2016). Thus, despite being efficient, these reproductive barriers may still be permeable, with putative hybrids observed in the field and in manual interspecific crosses.

In agreement with genetic data, our CVA clearly indicated morphological discontinuities (Fig. 6), supporting differentiation between the species. Morphological differentiation between V. simplex and V. scalaris is associated mainly with floral traits (scape length; length and width of the floral bracts; length and width of the bract floral scape; and pistil length) (Table 5). These results suggest that morphological traits between species can be involved in maintaining species boundaries; however, more studies are needed to confirm this hypothesis.

### Hybridization patterns across sympatric populations and reproductive barriers

The STRUCTURE results identified hybrids in all sympatric populations (Fig. 2) confirming previous hypotheses of hybridization between V. simplex and V. scalaris based on field observations and manual interspecific crosses (Neri et al. 2017).

Ours results revealed differences in genetic composition among the sympatric populations studied. The EBS
and RUS sympatric populations have higher numbers of hybrids (STRUCTURE analysis) than the RDB sympatric population. In addition, only in EBS and RUS populations were hybrids classified by NEWHYBRIDS, and they were mostly F2s, although some hybrids could not be classified (Fig. 3). The ability to identify and classify hybrid individuals through genetic analysis depends ultimately upon the number of diagnostic loci detected (Moccia et al. 2007), with different fixed alleles in each species. In this study, most loci were not diagnostic, probably due to the sharing of ancestral polymorphism between these incipient species. Similar results were found in another pair of Vriesea species, with difficult to distinguish hybrid classes (Zanella et al. 2016).

In this study, hybrid individuals did not present clear morphological distinctions, suggesting that most

Table 3. Characterization of populations of Vriesea simplex, V. scalaris and their hybrids, with 15 nuclear microsatellite markers, including the number of individuals sampled (N), number of alleles (A), number of private alleles (Ap), allelic richness (AR), variance in allele size (Var), observed (Hs) and expected (He) heterozygosity, inbreeding coefficient (Fis), and apparent outcrossing rate (ta) for each population. Inbreeding coefficient (Fis) departed significantly from HWE are indicated by asterisks (*P < 0.05).

| Species (samples size) | N  | A  | Ap | AR | Var | Hs | He | Fis | ta |
|------------------------|----|----|----|----|-----|----|----|-----|----|
| Allopatric—V. simplex   |    |    |    |    |     |    |    |     |    |
| Guapimirim, RJ         | 16 | 111| 18 | 6.75| 33.67| 0.545| 0.668| 0.196*| 0.693|
| Soberbo, RJ            | 28 | 157| 32 | 7.80| 37.14| 0.584| 0.732| 0.215*| 0.658|
| Allopatric—V. scalaris |    |    |    |    |     |    |    |     |    |
| Peri, SC               | 20 | 29 | 4  | 1.88| 17.19| 0.195| 0.221| 0.145*| 0.753|
| Sincorá, BA            | 10 | 33 | 7  | 2.20| 17.34| 0.134| 0.252| 0.509*| 0.408|
| Simpatric—Santa Lucia, ES |    |    |    |    |     |    |    |     |    |
| V. simplex             | 57 | 170| 46 | 6.50| 27.59| 0.556| 0.649| 0.178*| 0.704|
| Hybrid                 | 12 | 83 | 50 | 2.33| 29.09| 0.411| 0.594| 0.318*| 0.592|
| V. scalaris            | 39 | 55 | 19 | 2.73| 36.55| 0.105| 0.312| 0.673*| 0.158|
| Simpatric—Ruschi, ES   |    |    |    |    |     |    |    |     |    |
| V. simplex             | 18 | 102| 9  | 6.11| 42.04| 0.538| 0.649| 0.204*| 0.691|
| Hybrid                 | 8  | 64 | 20 | 2.26| 33.14| 0.359| 0.557| 0.371*| 0.653|
| V. scalaris            | 14 | 42 | 8  | 2.69| 16.26| 0.084| 0.385| 0.797*| 0.225|
| Simpatric—Rebio Duas Bocas, ES |    |    |    |    |     |    |    |     |    |
| V. simplex             | 16 | 86 | 6  | 5.71| 23.66| 0.545| 0.616| 0.102*| 0.845|
| Hybrid                 | 2  | 20 | 6  | 1.33| 28.85| 0.411| 0.594| 0.200*| 1.000|
| V. scalaris            | 15 | 34 | 11 | 2.16| 19.90| 0.187| 0.359| 0.435*| 0.358|

Table 4. AMOVA for 15 nuclear microsatellites with two hierarchical levels, including Vriesea simplex and V. scalaris pure individuals in sympatric and allopatric populations.

| Source of variation | Variation % | F-statistics | P-value |
|---------------------|-------------|--------------|---------|
| By species          |             | Fst =0.10999 | <0.001  |
| Among species       | 10.99       |              |         |
| Among population within species | 16.86 | Fst =0.19086 | <0.001  |
| Within populations  | 74.01       | Fst =0.27986 | <0.001  |
| Vriesea simplex     |             |              |         |
| Among populations   | 6.98        | Fst =0.069   | <0.001  |
| Within populations  | 93.01       |              | <0.001  |
| Vriesea scalaris    |             |              |         |
| Among populations   | 41.67       | Fst =0.416   | <0.001  |
| Within populations  | 58.32       |              | <0.001  |
hybrids may be identified as a parental species based only on morphology (Fig. 6). Although the hybrids are not intermediates, some individuals have an unusual morphology when compared to the parent species. In fact, it is well known that morphological traits alone are limited when identifying natural hybrids, especially considering incipient species (e.g. Li et al. 2015a, b; Moreno et al. 2015). Although hybrids with intermediate morphologies were not clearly observed, the occurrence of individuals with intermediate admixture values in all sympatric populations indicates that hybridization events are likely. In agreement with this, there is an overlap of blooming and pollinators (P. eurynome and R. naevious) in sympatric areas (Varassin and Sazima 2000; Wendt et al. 2008), which potentially favour interspecific pollen exchange. Thus, the overlap of flowering and pollinators can be considered as less effective prezygotic barriers in this system (Wendt et al. 2008).

However, in the face of hybridization, reproductive isolation may be maintained (Coyne and Orr 2004) and other prezygotic and postzygotic reproductive barriers can contribute to isolation between two species (Rieseberg and Willis 2007; Lowry 2008; Widmer et al. 2009).

Our analysis of two hybridizing Vriesea species allows us to discuss the general barriers involved in the maintenance of species integrity. Differences in the reproductive system of these species (Neri et al. 2017), with the predominance of selfing in V. scalaris and outcrossing in V. simplex, may be considered as a premating reproductive isolation barrier. We observed sympatric populations in V. scalaris with lower $t_s$ values (0.158–0.358) than in allopatric populations (0.408–0.754). This difference strongly suggests that in sympatry, V. scalaris tends to have higher selfing than in allopatry. Furthermore, we observed asymmetric levels of gene flow from V. scalaris into V. simplex (Fig. 5), suggesting selfing as a potential reproductive barrier between these species. In fact, empirical (Fishman and Wyatt 1999; Martin and Willis 2007; Matallana et al. 2010; Brys et al. 2010)
Species integrity is maintained in the occurrence of hybridization

(2013, 2015; Palma-Silva et al. 2015) and theoretical (Hu 2015) studies have shown selfing in a potentially interbreeding species can affect rates of interspecific gene flow to an outcrossing species, contributing to reproductive isolation.

Divergent mating systems were reported to contribute as reproductive barriers in other plant species (Fishman and Wyatt 1999; Lowe and Abbott 2004; Fishman and Stratton 2004; Martin and Willis 2007), including bromeliads (Wendt et al. 2002; Matallana et al. 2010; Palma-Silva et al. 2015; Wagner et al. 2015). The divergent mating systems and asymmetric levels of gene flow may be a consequence of higher herkogamy (the distance between the stigma and anthers) in *V. simplex* than in *V. scalaris* (Neri et al. 2017). Hermogamy in *V. simplex* may increase the possibility of contact with heterospecific pollen. In contrast, in *V. scalaris*, with lower herkogamy, spontaneous selfing can facilitate the protection of stigmas with plant self-pollen, and may counterbalance the input of cross-pollen.

Our morphometric data showed significant variation in floral traits (floral bract length and width, scape bract length and scape length) among sympatric and allopatric populations of *V. simplex* (Fig. 7). Floral bract length and width and scape bract length in *V. simplex* were larger in sympatric populations than in allopatric populations. These results suggest that sympatric floral display in the outcrosser *V. simplex* tends to be showier than in allopatry. In addition, we observed a higher effective migration rate (\(N_{e,m}\)) towards *V. simplex* in the sympatric populations, which reflects diversity and genetic structure, as well as the variation in flower traits of this species in the hybrid zones.

In contrast, no significant variations in floral traits were found among the sympatric or allopatric populations of *V. scalaris*. In fact, these floral phenotypes are often assumed to be the result of pollinator selection (Barrett and Harder 1996). The divergence of floral traits in sympatric and allopatric populations may evolve in response to competition for pollinator resources, or in response to the costs associated with pollinator sharing between species (Beans et al. 2014). Christianini et al. (2012) studying sympatric populations of the bromeliad genus *Encholirium* observed that divergence in floral traits and pollinator assemblage may contribute to reproductive isolation between species. Further investigation of the genetic basis of floral traits, including bract colour, in these *Vriesea* species and its interaction with pollination, will shed light on the specific role of floral display in reproductive isolation between closely related species. In *Mimulus* species, two genes altering flower colour were responsible for pollinator shifts and considered an important barrier to maintaining species boundaries (Ramsey et al. 2003; Yuan et al. 2013). Forthcoming studies on the fitness of sympatric

![Figure 6. Scatter plot of the scores derived from discriminant functions CVA1 versus CVA2 produced by stepwise discriminate analysis (CVA) applied to 24 morphometric characters for *Vriesea simplex* and *V. scalaris.*](image)

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populations versus allopatric populations may indicate whether these floral trait variations are due to reinforcement or to ecological character displacement.

In addition to the prezygotic reproductive barriers discussed above, the low germination rate of interspecific crosses observed in a previous study (Neri et al. 2017) suggest that postzygotic isolation may also be involved in maintaining reproductive isolation between *V. simplex* and *V. scalaris*. Lower hybrid seed viability could also explain the low frequency of hybrids observed in nature. Reduced seed viability in interspecific crosses may be due to genetic incompatibility, as in BDM incompatibility (Orr and Turelli 2001; Welch 2004), resulting from negative genetic interaction among nuclear-nuclear loci (Orr and Turelli 2001; Bomblies et al. 2007) or cytoplasmic-nuclear loci (Greiner et al. 2011). Inviable or sterile hybrids due to genetic incompatibility are potential postzygotic barriers preventing parental species collapse in hybrid zones (Coyne and Orr 2004; Scopece et al. 2010). The accumulation of genetic incompatibility was also observed in other pairs of plant species (Moyle and Nakazato 2010; Scopece et al. 2008, 2010; Palma-Silva et al. 2011; Moyle et al. 2012; Ishizaki et al. 2013; Briscoe Runquist et al. 2014; Johnson et al. 2015; Pinheiro et al. 2015; Matallana et al. 2016).

**Conclusions and Prospects**

Here we show that genetic and morphological integrity between *V. simplex* and *V. scalaris* are maintained despite natural hybridization. Our data suggest that in sympatric populations *V. scalaris* tends to have higher selfing rates than in allopatric populations, suggesting that selfing can potentially reduce rates of interspecific gene flow from an outcrossing species. Complementary isolating mechanisms, such as variation in floral traits, among sympatric and allopatric populations in the outcrosser *V. simplex*, may also contribute to the maintenance of species integrity, due to stronger floral display in sympatric populations. The presence of multiple prezygotic and postzygotic barriers and their interactions, although still permeable, probably allow these species to persist in sympatry. While flowering time and pollinator specificities do not appear to be effective prezygotic barriers, we observed that the reproductive system, including floral traits and low seed viability, might contribute to species integrity. To obtain a more complete picture of the species composition of a hybrid zone, it will be necessary in future studies to use a combination of morphological characters and a larger genomic data set that combines nuclear and plastidial markers.

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**Contributions by the Authors**

J.N. conceived the ideas/conducted the collections, conducted experiments, performed the work in the laboratory/analysed data and led the writing of the manuscript.

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**Table 5.** Standardized coefficients for canonical variables derived from discriminant function analysis (CVA) from 11 populations of *Vriesea simplex* and *V. scalaris* along the Atlantic rainforest, Brazil. The sublimated characters are those that contribute the most to the species separation according to the values of CVA1 and CVA2.

| Characters             | CVA1     | CVA2     |
|------------------------|----------|----------|
| Scape length           | 0.407671 | 0.420156 |
| Floral bract width     | 0.540610 | −0.063002|
| Floral bract length    | 0.490502 | −0.293909|
| Scape bracts width     | 0.416668 | 0.118613 |
| Scape bracts length    | 0.362053 | −0.092482|
| Length pistil          | 0.332938 | 0.143569 |
| Leaf heath length      | 0.012159 | 0.050331 |
| Leaf heath width       | 0.177408 | 0.320468 |
| Leaf blade length      | 0.20957  | 0.610379 |
| Leaf blade width       | 0.203816 | 0.071918 |
| Inflorescence total length | 0.275906 | 0.382661 |
| Rachis length          | 0.125729 | 0.276047 |
| Flowers number         | 0.153994 | 0.101936 |
| Sepal width            | 0.128110 | 0.003864 |
| Sepal length           | 0.041543 | 0.062018 |
| Petal width            | 0.290863 | −0.054088|
| Petal length           | 0.005314 | 0.042810 |
| Anther length          | 0.077106 | −0.028683|
| Pedicle length         | 0.224256 | 0.022448 |
| Anther–stigma distance | 0.164429 | 0.026982 |
| Stamen length          | 0.294832 | 0.056851 |
| Fillet length          | 0.255827 | 0.090300 |
| Rosette diameter       | 0.151645 | 0.005057 |
| Rosette height         | 0.243857 | 0.150933 |
C.P.S. conceived the ideas/conducted the collections/contributed with reagents/materials/analysed data and led the writing of the manuscript. T.W. conceived the ideas/collaborated with materials/revised the writing. This manuscript is part of the PhD thesis of the first author.

Conflict of Interest
None declared.

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Supporting Information
The following additional information is available in the online version of this article—
Table S1. Voucher of the populations collected in this study.
Table S2. Summary of the morphometrics of Vriesea simplex, V. scalaris and their hybrids.
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