Patterns of style polymorphism in five species of the South African genus *Nivenia* (Iridaceae)

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INTRODUCTION

Discrete sexual polymorphisms in plants, i.e. the existence of two or more morphs affecting sexual organs within single populations, is a pervasive trait in angiosperms which has attracted evolutionary biologists ever since Darwin’s seminal work (Darwin, 1877). The species having these genetically based polymorphisms are excellent model systems to study the functioning and evolution of the mechanisms for disassortative mating. Among these polymorphisms, heterostyly is by far the most common. Its functional significance is thought to be simultaneously the avoidance of self-interference of sexual whorls between morphs [although Lloyd and Webb (1992) provide an intermediate stage of stigma-height dimorphism (two morphs with different style lengths but the anthers at the same level). Stigma-height dimorphism is considered to be unstable because of lack of precision in pollen transfer between morphs [although Lloyd and Webb (1992) propose that the ancestral condition is an approach herkogamous flower (stigmas above anthers, the commonest arrangement in angiosperms; Webb and Lloyd, 1986), then an intermediate stage of stigma-height dimorphism (two morphs with different style lengths but the anthers at the same level). Stigma-height dimorphism is considered to be unstable because of lack of precision in pollen transfer between morphs [although Lloyd and Webb (1992b) propose specific conditions for its maintenance; for a test case, see Cesaro and Thompson, 2004], thus a reciprocal positioning of sexual whorls between morphs would be selected for. The selective factor is therefore the promotion of the efficiency in pollen transfer driven by the particular behaviour of the pollinators collecting and delivering pollen grains in precise parts of the flower sexual whorls and some traits considered as ancillary are supplied to determine for each population (a) the kind of stylar polymorphism, (b) the morph ratio and (c) the degree of reciprocity between sexual whorls. Also the rates of assortative (within morph) versus disassortative (between morphs) pollen transfer were estimated by analysing pollen loads on stigmas. The association between floral phenotypic integration and the reciprocity between sexual whorls was estimated; both characteristics have been quoted as dependent on the accuracy of the fit between pollinators and flowers and therefore related to the efficiency of pollen transfer.

Key Results Different types of polymorphism, differing in their degree of reciprocity, were found in *Nivenia*. Effective disassortative mating appears to be common, since (a) all dimorphic populations show equal morph-ratios (isoplethy), and (b) the pollen placed on the stigmas of each morph is likely to be coming from the other (complementary) morph. The most reciprocal populations of the heterostylous species have also the highest values of phenotypical integration.

Conclusions Stigma height dimorphism, as opposed to distyly, is proven for the first time in *Nivenia*. The presence of different types of polymorphism within the genus is consistent with hypotheses of the evolution of heterostyly. The role of the pollinators as the leading force of the transition seems to be apparent, since floral integration is related to reciprocity.

Key words: Heterostyly, *Nivenia*, phenotypic integration, reciprocal herkogamy, reciprocity degree, disassortative mating.
of their bodies. Usually heterostyloous plants also have other typical features, such as a diallelic incompatibility system which only permits crosses between morphs (for a review, see Barrett and Cruzan, 1994), and some between-morph differences in floral traits (ancillary traits: e.g. pollen and stigma size and shape, corolla size, etc.; Ganders, 1979). In Lloyd and Webb’s (1992a) model, these features are interpreted as devices to increase the efficiency of the system by avoiding gamete wastage. Thus incompatibility would appear after reciprocal herkogamy, both traits not necessarily being linked.

In recent times the depicted evolutionary scenario has gained increasing support from diverse areas such as pollination ecology, flower ontogeny, breeding systems and phylogenetics. Nevertheless, despite the wide taxonomic representation of heterostyly among angiosperms [28 families (Barrett, 1992) and 164 genera (Ganders, 1979)], most studies have somewhat concentrated on only a few taxa [e.g. Pontederiaceae (Barrett et al., 1989), Turneraeae (Shore et al., 2006), Primulaceae (Nishihiro et al., 2000), Rubiaceae (Fairev and McDade, 2001), Oxalidaceae (Weller et al., 2007), Amarillidaceae (Barrett and Harder, 2005; Pérez-Barralés et al., 2006), Boraginaceae (Schoen et al., 1997; Brys et al., 2008a, b; Ferrero et al., 2009)]. For example, stigma-height dimorphism (as opposed to distyly, sensu Barrett et al., 2000) has only recently been shown to be the most likely intermediate step in Narcissus (Graham and Barrett 2004; Pérez-Barralés et al., 2006) and Lithodora sensu lato (Ferrero et al., 2009). Although stigma-height dimorphism has been reported in other groups where heterostyly does exist [Anchusa (Philipp and Schou, 1981), Quichamalium (Riveros et al., 1987)], the evolutionary relationship between distyly and dimorphism remains to be proven. Since stigma-height dimorphism has sometimes mistakenly taken as distyly (e.g. Fernandes, 1964; Philipp and Schou, 1981; Riveros et al., 1987; Richards and Koptur, 1993) stigma-height dimorphism is likely to be more frequent than was previously thought, and therefore distyly overestimated. Detailed floral morphometric studies from different groups where heterostyly and monomorphism are present are thus necessary. Such morphometric studies would throw light on at least two complementary aspects: (1) measurements of sexual organs could show the presence of the supposed intermediate stages, and are needed to estimate the contribution of each variant to fitness; (2) perianth traits could reveal the importance of precise pollinators, which may be inferred from restrictive corollas (Lloyd and Webb, 1992a) and strong correlations among floral traits (Berg, 1960). In fact, higher correlations among floral traits (phenotypic integration sensu Wagner, 1984) were found in style polymorphic populations than in monomorphic populations (Pérez-Barralés et al., 2007). Moreover, a positive correlation between reciprocity and floral phenotypic integration has been found (Ferrero et al., 2011).

Nivenia is the only genus within the Iridaceae containing heterostyloous species, and has been reported to include distyloous and monomorphic species (Mulcahy, 1965; Ornduff, 1974; Goldblatt, 1993). The taxonomic monograph by Goldblatt (1993) reported four monomorphic species and five heterostyloous species. Afterwards Goldblatt (1997) described N. parviflora, and recently Manning and Goldblatt (2007) have described a new species, N. inaequalis, both considered to be distyloous. The most comprehensive study on the floral biology of Nivenia is that of Goldblatt and Bernhardt (1990) which reports data on flower morphology, breeding systems, and pollinators for all the distyloous species known up to that date. Unfortunately, absolute measurements of sexual organs were not included there, and the relative lengths provided do not allow to determine if the considered distyloous species are truly so, nor their degree of reciprocity, although some apparent variability deserves further investigation. Nor has the incompatibility system been conclusively studied for the genus, contrasting the results of Goldblatt and Bernhardt (1990) that suggest at least partial morph-compatibility based on the growth of pollen tubes in the pistils, with those by Ornduff (1983) who reported morph-incompatibility for N. corymbosa based on seed production.

Here, variation in floral traits related to style polymorphism and pollination biology is reported in five species of Nivenia, to explore if this variation meets the requirements of any model for the evolution of heterostyly. A definite test would need to reconstruct evolutionary transitions of these variations on the phylogeny, which is not yet available.

The specific aims of this work are (a) to analyse the variation of floral traits in order to quantitatively characterize the type of floral polymorphism and its reciprocity level; (b) to determine the morph ratio in the populations; (c) to determine the pollen load on stigmas and to estimate the pollen transfer rate between morphs, and hence the efficiency of style polymorphism, using the between-morph variability in pollen size; and (d) to estimate the levels of floral phenotypic integration and its relationship with the polymorphism and the pollinator types, as well as the possible role of any particular floral trait in pollination success.

**MATERIALS AND METHODS**

**Study species and population sampling**

The genus Nivenia Vent. (Iridaceae, subfam. Nivenioideae) is endemic to the Western Cape Province (South Africa), and all species are restricted to sand-stones of the montane layer of the Cape System, on soils of the Table Mountain Series (Goldblatt and Bernhardt, 1990; Goldblatt, 1993). It comprises small to tall evergreen shrubs with woody underground stems. Inflorescences are terminal, forming a pseudopanicle with short to long axillary branches, thus becoming corymbose or raceme-like. Flowers, from one to many per inflorescence, are blue with well-developed perianth tubes and outspread tepals. The filaments of the stamens (three per flower) are adnate to the tube resulting from the connation of the six tepals. All three anthers of the flower are placed at the same height level except for N. inaequalis, which takes its name from this trait. The ovary is inferior, globose and usually two-ovulate (for a thorough description, see Goldblatt, 1993; Manning and Goldblatt, 2007).

Six populations of five species were surveyed between 1998 and 2005 (Table 1). Given the narrow endemicity of most species, sampling was non-destructive at the plant level; in

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each population, only one apparently healthy recently opened flower per plant was collected in 100 randomly selected plants when available (just one flower from dense clumps to avoid oversampling of genets), assuming a similar age for the collected flowers. Flowers were preserved in 70 % ethanol until their measurement in the laboratory. In addition, one flower-bud per plant was also sampled in 50 plants to estimate the size and production of pollen grains. According to our experience, that sampling number is representative of the populations, with a number in the population of a few hundreds at most; note that in populations with reduced population size, like *N. argentea* at Aasvoëlkrans, all flowering individuals were sampled.

**Morphometric measurements and morph ratios**

To characterize and evaluate the degree of reciprocity in each population, flowers were photographed, longitudinally dissected, and then photographed again. Measurements were taken from the digital photos with the image analyser software analySIS 5.0. Floral traits measured were: (a) stamens height (up to the insertion of the filament on anther for each one of them); (b) stigma height; (c) corolla tube length; (d) tube width; (e) tepal limb length; (f) bract length; and (g) anther length (Fig. 1). Sample sizes ranged between 10 and 100 (Table 1). Individual flowers were classed as L (long) morphs when the stigma level was above the level of the anthers, or S (short) morphs when the stigma level was below the level of the anthers.

To characterize each population as style-dimorphic (all stamens of both morphs at the same height level) or truly heterostylos (different stamen level for each morph), *t*-tests were performed to compare the heights of anthers between morphs. When anther heights do not show significant between-morph differences the population was considered as style-dimorphic. Also the fit between anther and stigma heights for each morph was calculated with regression. The regression coefficients were compared to find out if both sexual whorls co-varied at the same rate in both morphs.

Also the degree of reciprocity between sexual whorls for each population was calculated with the index of Sánchez et al. (2008). Unlike previous indices, it provides meaningful values that can be compared across populations and species because it compares stigma–stamen height gaps for all potential legitimate crosses (i.e. each and every stamen of a morph versus each and every stigma of the complementary morph measured in a population) while considering also the dispersion of the data, and it is not skewed by the more frequent sex (stamens). Unlike previous indices based only on a measure for stamen and stigma heights, this index needs a lot of computation to deal with all the available data of a population (for a complete description, see Sánchez et al., 2008; computational software available at [http://webs.uvigo.es/plantecology/software.es.html](http://webs.uvigo.es/plantecology/software.es.html)).

Among the studied ancillary traits were (a) the total number of pollen grains per anther and (b) pollen grain size. To this end, each anther was placed in a drop of 50 % glycerine over a microscope slide, opened and squashed beneath a cover slip; the pollen grains were counted under a light microscope (magnification × 100) differentiating among healthy and apparently aborted (collapsed) grains. The pollen preparations were photographed under the optical microscope and measurements of polar axis in 50 grains were made with the image analyser software analySIS 5.0.

Differences between morphs in each population were compared with a *t*-test for all traits, except when the available number of flowers was small (*N. binata* and *N. argentea* at Aasvoëlkrans) which were analysed with the non-parametric Mann–Whitney test. The numbers of aborted pollen grains were compared between morphs with ANCOVA, considering also the dispersion of the data, and it is not skewed by the more frequent sex (stamens). Unlike previous indices based only on a measure for stamen and stigma heights, this index needs a lot of computation to deal with all the available data of a population (for a complete description, see Sánchez et al., 2008; computational software available at [http://webs.uvigo.es/plantecology/software.es.html](http://webs.uvigo.es/plantecology/software.es.html)).

**FIG. 1.** Main floral parts measured in the flowers sampled of both morphs (L and S): (1) height of all stamens, (2) stigma height, (3) corolla tube length, (4) tube width, (5) length of the tepal limb, (6) bract length. Anther lengths were also measured but are not indicated in the figure due to small size (schematic drawing modified from Goldblatt, 1993).

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**TABLE 1. Summarized data of the localities of the five species of Nivenia studied**

<table>
<thead>
<tr>
<th>Species</th>
<th>Location</th>
<th>Altitude (m)</th>
<th>No. of plants sampled (L/S morphs)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>N. inaequalis</em></td>
<td>Rooiberg Massif, near Teeboskop peak, slope facing south (33° 41’S, 21° 34’E)</td>
<td>1021</td>
<td>100 (53/47)</td>
</tr>
<tr>
<td><em>N. argentea</em></td>
<td>Phesantefontein, near Aasvoëlkrans peak, slope facing south (33° 56’S, 21° 08’E)</td>
<td>955</td>
<td>10 (6/4)</td>
</tr>
<tr>
<td><em>N. argentea</em></td>
<td>Peak south Langkloof, near Garcia pass, crest and slope south (33° 57’S, 21° 15’E)</td>
<td>1033</td>
<td>100 (59/41)</td>
</tr>
<tr>
<td><em>N. binata</em></td>
<td>Swartberg Pass (33° 21’S, 22° 03’E)</td>
<td>1228</td>
<td>18 (13/5)</td>
</tr>
<tr>
<td><em>N. corymbosa</em></td>
<td>Wellington, Bain’s Kloof Pass (33° 38’S, 19° 05’E)</td>
<td>451</td>
<td>84 (35/52)</td>
</tr>
<tr>
<td><em>N. fruticosa</em></td>
<td>Perdeberg (33° 56’S, 21° 04’E)</td>
<td>1144</td>
<td>18</td>
</tr>
</tbody>
</table>

To estimate the morph ratio in each polymorphic population. Deviations from isoplethy (equal morph-ratio) were tested by means of the chi-square test.
Pollen transfer within populations

Between-morph differences in pollen reception were tested by comparing pollen loads on stigmas. According to Goldblatt and Bernhardt (1990) and the present observations, pollen size dimorphism in the polymorphic species of Nivenia could allow to discrimination between the pollen produced by each morph. Based on this possibility, the polar axis of pollen grains on stigmas was measured with the image analyser software analySIS 5.0, to assess the level of assortative and disassortative pollen flow within the populations.

Hence, styles were cleared and softened with 8 N sodium hydroxide for 24 h, rinsed in distilled water and stained overnight with 0.05% aniline blue prepared in 0.1 M potassium phosphate (Dafni et al., 2005). Pistils were then placed on a microscope slide with a drop of 50% glycerine and squashed beneath a cover slip. Samples were observed and photographed through a Nikon Eclipse 80i epifluorescence microscope (Nikon Instruments, Kanagawa, Japan) with a UV-2A filter cube and the following variables were measured on these photographs: (a) number of pollen grains on the stigma; (b) polar axis diameter of five, when possible, pollen grains randomly chosen in each stigma; and (c) number of pollen tubes inside the style. Pollen loads on stigmas were estimated as the addition of the number of pollen grains on the stigma plus the number of pollen tubes in the style when not linked to a pollen grain.

To compare pollen reception between morphs 2 × 2 contingency tables for each population were analysed, considering the morph (long/short styled) and presence/absence of pollen on the stigma. Since the size of the pollen ranges overlapped between morphs (and therefore it cannot be conclusively decided by which morph a particular pollen grain was produced), the prevalence of assortative or disassortative mating was assessed by comparing (t-test) the mean sizes of the pollen grains on the L and S stigmas.

RESULTS

Morphometric analysis and morph ratio

Average values of the measurements of the floral traits for each population and index of reciprocity between sexual whorls are shown in Tables 2 and 3, respectively. Variation of sexual whorls is represented in comparative plots in Figs 2–5. Among the species studied, only N. fruticosa is monomorphic, presenting clearly approach herkogamous flowers where the style is located well above the anthers (Table 2). For the species with two floral morphs, stigma height was significantly different in all of them (Table 2), and the length of the stamens was statistically different between morphs in all cases, except for the population of N. argentea at Aasvoelkrans (Table 2). Thus, those were considered as distylous, and N. argentea at Aasvoelkrans as stigma-height dimorphic since two style-length morphs are present but anther heights remain indistinguishable between morphs (see Fig. 3A). It has been thus confirmed that this is not a biased sample from a larger distylous population, but a truly stigma-height dimorphic, with a bootstrapping simulation on the data of the population of N. argentea at Garcia (distylous, n = 100). Re-sampled subpopulations with the same sample size of the population at Aasvoelkrans (n = 10, 6 large- + 4 short-styled flowers) were randomly extracted, and the mean stamen height compared between morphs at each iteration (t-test, P < 0.01) in MsExcel®. Since the probability of obtaining stigma-height dimorphic subsamples from the large heterostylous population at Garcia is negligible (1000 iterations, P < 0.0001), it can be concluded that the population at Aasvoelkrans is truly stigma-height dimorphic.

The results show that N. inaequalis has higher and more dispersed stamens and stigmas than N. argentea (Figs 2 and 3). A larger dispersion is in part due to the fact that in N. inaequalis one of the three stamens is clearly placed under the level of the other two. When comparing stigma and mean stamen height for N. inaequalis (Fig. 2B) two facts can be readily noticed: (1) the high dispersion of the values; (2) their fitted lines are close to parallel (i.e. their regression coefficients are not statistically different, t = 1.92, P = 0.06), meaning that the height of stamens and stigmas varies isometrically in both morphs. For each population two values were calculated, one for each morph.
<table>
<thead>
<tr>
<th>N. inaequalis</th>
<th>N. argentea Aasvoëlkrans</th>
<th>N. argentea Garcia</th>
<th>N. binata</th>
<th>N. corymbosa</th>
<th>N. fruticosa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Morph</td>
<td>n</td>
<td>Mean ± s.d.</td>
<td>n</td>
<td>Mean ± s.d.</td>
<td>n</td>
</tr>
<tr>
<td>Floral characteristics (mm)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stamen height</td>
<td>L</td>
<td>158</td>
<td>35.25 ± 4.74**</td>
<td>18</td>
<td>22.09 ± 1.64</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>138</td>
<td>37.28 ± 3.84**</td>
<td>12</td>
<td>21.93 ± 1.18</td>
</tr>
<tr>
<td>Stigma height</td>
<td>L</td>
<td>53</td>
<td>41.38 ± 2.95**</td>
<td>6</td>
<td>27.19 ± 1.84**</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>47</td>
<td>30.90 ± 2.80**</td>
<td>4</td>
<td>13.62 ± 1.55**</td>
</tr>
<tr>
<td>Tube diameter</td>
<td>L</td>
<td>53</td>
<td>02.80 ± 0.32</td>
<td>6</td>
<td>03.34 ± 0.44</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>47</td>
<td>02.81 ± 0.33</td>
<td>4</td>
<td>03.29 ± 0.34</td>
</tr>
<tr>
<td>Tube length</td>
<td>L</td>
<td>53</td>
<td>33.63 ± 3.31**</td>
<td>6</td>
<td>20.98 ± 1.62*</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>47</td>
<td>32.28 ± 2.93**</td>
<td>4</td>
<td>17.90 ± 0.98*</td>
</tr>
<tr>
<td>Tepal limb length</td>
<td>L</td>
<td>53</td>
<td>13.40 ± 1.66</td>
<td>6</td>
<td>11.11 ± 1.03</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>47</td>
<td>13.47 ± 1.50</td>
<td>4</td>
<td>11.28 ± 0.89</td>
</tr>
<tr>
<td>Bract length</td>
<td>L</td>
<td>53</td>
<td>23.56 ± 2.38</td>
<td>6</td>
<td>20.76 ± 2.95</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>47</td>
<td>22.83 ± 2.85</td>
<td>4</td>
<td>17.59 ± 3.40</td>
</tr>
<tr>
<td>Anther length</td>
<td>L</td>
<td>133</td>
<td>01.73 ± 0.24</td>
<td>18</td>
<td>01.88 ± 0.18**</td>
</tr>
<tr>
<td></td>
<td>S</td>
<td>65</td>
<td>01.80 ± 0.21</td>
<td>12</td>
<td>02.25 ± 0.20**</td>
</tr>
</tbody>
</table>

Pollen characteristics in buds [n = number of anthers (number of flowers)]

|                      | L | 21 (7) | 874.1 ± 184.9 | 15 (5) | 2169.2 ± 322.4 | 21 (7) | 2083.5 ± 209.1 | – | – | 21 (7) | 2236.7 ± 405.5 | 30 (10) | 787.3 ± 246.2 |
|                      | S | 21 (7) | 917.0 ± 228.1 | 12 (4) | 2383.8 ± 275.4 | 21 (7) | 2199.1 ± 360.6 | – | – | 21 (7) | 2295.8 ± 387.3 | 30 (10) | 142.5 ± 310.5 |

Aborted pollen grains per anther [n: number of grains (number of flowers)]

|                      | L | 21 (7) | 121.2 ± 10.1 | 15 (5) | 46.2 ± 44.5 | 21 (7) | 111.8 ± 121.7 | – | – | 21 (7) | 125.5 ± 219.1 | 30 (10) | 142.5 ± 310.5 |
|                      | S | 21 (7) | 20.7 ± 21.5 | 12 (4) | 32.8 ± 15.1 | 21 (7) | 90.1 ± 74.3 | – | – | 21 (7) | 29.2 ± 20.7 | – | – |

Pollen grain size (µm)

|                      | L | 525 (7) | 53.35 ± 4.63** | 375 (5) | 45.81 ± 4.77** | 525 (7) | 42.41 ± 4.19** | 225 (7) | 42.17 ± 4.69** | 525 (7) | 39.81 ± 4.00** | 750 (10) | 55.34 ± 7.05 |
|                      | S | 525 (7) | 54.67 ± 4.92** | 300 (4) | 44.25 ± 4.45** | 525 (7) | 46.82 ± 4.39** | 75 (7) | 46.16 ± 3.66** | 525 (7) | 40.75 ± 4.48** |

Approach herkogamous flowers were categorized as L, and reverse herkogamous as S. Morphs within populations were compared with a t-test, except that (a) comparisons within the populations of N. binata and N. argentea were performed with non-parametric test (Mann–Whitney) due the small sample size; (b) the number of aborted pollen grains was compared with ANCOVA considering the total number of grains per anther as a co-variable, except for N. binata and N. argentea, where the proportion of aborted pollen grains was compared with non-parametric test (Mann–Whitney).

Values in bold differ significantly between morphs (* P < 0.05, ** P < 0.001).
This species presents high between-morph reciprocity, that is, a low $R$ index (Table 3).

The two populations of *N. argentea* (Aasvoëlkraans and Garcia) are represented in Fig. 3. As previously stated, the population at Aasvoëlkraans is stigma-height dimorphic; while the population at Garcia is distylous, but the stigmas of the short-styled morph seem to be lacking reciprocal stamens, since the anthers of the long-styled morph are placed constantly above the level of the short-styled stigmas (Fig. 3B). The relationship between stigmas and stamens at the intra-flower level for the population with a large sample size (Garcia, Fig. 3C) shows a diverging pattern between the fitted lines of the morphs (i.e. statistically significant difference between the regression coefficients, $t = 2.45$, $P = 0.02$), reflecting an allometric behaviour of the sexual organ heights between morphs. This pattern reflects certain independence between the ontogenetic growth of stigmas and stamens, especially for the short morph where the fitted line is closer to horizontal. The reciprocity degree for this species is the lowest (the highest $R$-values) of the four species studied (Table 3).

*Nivenia binata* is distylous with a small separation between the heights of stamens of the short and long morphs (Fig. 4A), and a high reciprocity degree (low $R$-values; Table 3). The relationship between stigma and mean stamen heights for

### Table 3. Reciprocity values ($R$) after Sánchez et al. (2008) (note that the higher the value, the lower the reciprocity); floral phenotypic integration for each morph as a percentage of the maximum possible value (number of traits considered); proportion of pollinated flowers; and pollen load range and median in pollinated flowers, for the style polymorphic populations

<table>
<thead>
<tr>
<th>Population</th>
<th>$R$</th>
<th>Phenotypic integration (%)</th>
<th>No. of flowers pollinated/unpollinated (%)</th>
<th>Pollen load ranges in pollinated flowers [min–max (median)]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>L</td>
<td>S</td>
<td></td>
</tr>
<tr>
<td><em>N. inaequalis</em></td>
<td>0.019</td>
<td>44.1</td>
<td>48.7</td>
<td>44/56* (44-0)</td>
</tr>
<tr>
<td><em>N. argentea</em> Aasvoëlkraans</td>
<td>0.043</td>
<td></td>
<td></td>
<td>6/10* (60-0)</td>
</tr>
<tr>
<td><em>N. argentea</em> Garcia</td>
<td>0.031</td>
<td>47.0</td>
<td>32.9</td>
<td>65/35** (65-0)</td>
</tr>
<tr>
<td><em>N. binata</em></td>
<td>0.021</td>
<td></td>
<td></td>
<td>7/19** (38-9)</td>
</tr>
<tr>
<td><em>N. corymbosa</em></td>
<td>0.019</td>
<td>52.2</td>
<td>51.7</td>
<td>50/38** (56-8)</td>
</tr>
</tbody>
</table>

Phenotypic integration was not calculated for populations with a sample size which was too small.

Differences between the number of pollinated and unpollinated flowers were tested with chi-square (n.s., not significant; **P < 0.001).
diverging fitted lines emerge, suggesting an allometric height for each flower (Fig. 5B), two clearly addition, when stigma height is compared with the mean Reciprocity between morphs (the lowest separated sexual whorls (Fig. 5A) and the highest degree of morphs (Fig. 4B).

_N. corymbosa_ flowers of \( N. inaequalis \) and \( N. argentea \) have larger tubes in the short-morph, while the latter has significantly shorter tubes in the short-morph. Tepal limb length varies between morphs but only in the population of \( N. argentea \) at Garcia (Table 2).

Anthers and pollen grains are slightly bigger in the short-styled flowers for all species (Table 2). In relation to pollen grain production, the anthers of both morphs produce a similar amount of pollen grains in each population, either healthy or aborted (Table 2).

The morph ratio of all polymorphic populations was isoplethic, i.e. 50 % of each morph in the population (\( \chi^2 \) test, \( P \) ranging between 0.006 (\( N. binata \)) and 0.55 (\( N. inaequalis \)).

### Pollen transfer within populations

Pollen transfer in the distylous populations of _Nivenia_ analysed is mainly disassortative: pollen grains on stigmas of the long-styled flowers are on average larger and on the short-styled flowers are smaller (Fig. 6), although the difference is statistically significant only for _N. corymbosa_ (\( t = 2.90 \), d.f. = 156, \( P = 0.004 \)).

At the population level, no correlation has been found between reciprocity and the two components of the female fitness studied, i.e. number of flowers pollinated or mean pollen load found on stigmas of pollinated flowers (\( r_s = 0.1 \), \( r_s = 0.2 \), respectively, \( P > 0.05 \); Table 3).

The results showed a relationship between morph and presence/absence of pollen only for _N. inaequalis_, where long-styled flowers seemed to be more efficient in capturing pollen. Significant differences were not found in any of the other species studied (Table 4).

### Phenotypic integration

The average value of flower phenotypic integration for the populations was 46.7 %. Variation among populations, species and morphs was high (range: 32.9 % in the short-styled morph of _N. argentea_ at Garcia to 52.2 % in the long-styled morph of _N. corymbosa_). Morphs have similar values in _N. inaequalis_ and _N. corymbosa_, but short-styled morph of _N. argentea_ at Garcia (Table 3). The monomorphic _N. fruticosa_ presented high integration values (50.1 %), close to the values of _N. corymbosa_.

### Relationship between individual floral traits and pollination success

The results of the logistic regression analyses show that there is only a slight influence on the probability of having...
pollen on stigmas for the following cases: 'stigma height' of
the long-styled flowers of N. inaequalis \( (R^2 = 0.30, B =
2.07, P = 0.05) \), 'corolla tube diameter' for the long morph of
N. argentea \( (R^2 = 0.19, B = 1.11, P = 0.02) \), and 'corolla tube length' for the
short morph of N. corymbosa \( (R^2 = 0.19, B = 1.11, P = 0.02) \). No significant effect was found for the
remaining floral traits analysed.

**DISCUSSION**

**Morphological variation**

The comprehensive model explaining the evolution of hetero-
styly of Lloyd and Webb (1992a) assumes a transition from
of the six unsampled species are approach herkogamous monomorphic according to taxonomic descriptions (Goldblatt, 1993), like the one here confirmed as monomorphic based on detailed measurements (N. fruticosa). Interestingly, among the four polymorphic species sampled, variation was found in the patterns of polymorphism, both at population and species levels, as a result of our detailed morphometric analysis. Styolar polymorphism is by definition a population trait, and taxonomic information based on limited sampling within populations may produce spurious results or ignore small but meaningful variations (see, for example, Barrett et al., 1996, 1997; Armbuster et al., 2006; Sánchez et al., 2008).

Broadly, the present data fit the morphometric description of these species provided by Goldblatt (1993) for all floral traits except in the case of N. argentea. This is not surprising since Goldblatt (1993) characterized N. argentea with data of specimens from the population at Rooiberg (also studied here, Table 1), but this population has been recently described as the new species N. inaequalis by Manning and Goldblatt (2007), splitting it off N. argentea mostly because the inequality of the stamens of the former. Therefore, the measurements of the flowers of both N. argentea and N. inaequalis, taken in the present study, fit the description of Manning and Goldblatt (2007). Nivenia argentea shows a greater variation since the two populations studied have different types of polymorphism, namely distylly at Garcia and stigma-height dimorphism at Aasvoelkrans. As far as is known this is the first report of such variation within a single species, although it has been reported within some genera, e.g. Narcissus (Barrett et al., 1996) and Lithodora (Ferrero et al., 2009). It could be argued that the stigma-height dimorphism of the Aasvoelkrans population is the result of their small population size, and that the array of phenotypes is a biased ‘sample’ of the general pattern of the species; in such a scenario stigmalheight dimorphism would be a spurious pattern resulting from the random colonization by long-styled flowers of relatively higher long stamens and short styled flowers of shorter stamens (see Fig. 3B for the distylous distribution at Garcia). However, the probability of this randomly driven fixation has been discarded with the re-sampling simulation used: it can be argued that the stigmal-height dimorphism of the Aasvoelkrans population is real and not an artefact. Moreover, this population, though small, shows an isoplethic morph ratio; both factors (dimorphism and isoplethy) seem to indicate that this population has achieved high levels of disassortative mating. It would be worth, however, exploring more populations to ascertain the extent of the variation within N. argentea.

The lack of differences in the ancillary floral traits except pollen size (Table 2) supports the idea that they are not so critically tightened to the sexual polymorphism, as has been found in many studies (Ganders, 1979; Dulberger, 1992).

Functional aspects

Reciprocity values turned out to be the highest, and equal, in distylous N. corymbosa and N. inaequalis. Close values of reciprocity have been obtained for the distylous N. binata. The two populations of N. argentea were clearly less reciprocal than N. inaequalis (Table 3). Surprisingly, the higher degree of reciprocity of N. inaequalis is largely due to the higher dispersion of its anther heights (see Fig. 2A), given the displacement of the shortest stamen with respect to the level of the other two (Manning and Goldblatt, 2007; and this study). For this species, it was checked (and discarded) that the value of reciprocity was different when each anther height was considered separately: if that were the case, some anthers could have different probability of between-morph outcrossing, and therefore a different performance as male. In contrast, all three anthers of N. argentea flowers are placed at the same level. The greater dispersion of the anthers in N. inaequalis makes it possible that all stigmas have at least some reciprocal anthers (i.e. anthers of the other morph) at its level. This distinctly different arrangement of the androecium of N. inaequalis within the genus could be the result of directional selection on the length of one of the stamens in order to increase the probability of delivering pollen to legitimate stigmas. A similar process has been described for Lithodora by Ferrero et al. (2009). Both morphs are equally good in receiving pollen, as in other truly distylous species (N. binata and N. corymbosa; Table 3).

The greater ontogenetic independence of styles and stamens in N. corymbosa and N. argentea at Garcia is noteworthy. It is hypothesized that such independence allows the sexual organs to respond differently to the selective pressures imposed by the pollinators, which is the main force driving the evolution of heterostyly according to the Lloyd and Webb (1992a) model. Indeed such a scenario could have driven N. corymbosa to higher values of reciprocity than the other distylous species (Table 3).

Pollen transfer within populations

Although the presence of a diallelic incompatibility system may determine by itself an isoplethic morph ratio (Charlesworth and Charlesworth, 1979; Heuch, 1979), the available data for Nivenia (Goldblatt and Bernhardt, 1990) are not conclusive. There is increasing evidence that heterostyly and heteromorphic incompatibility are not necessarily linked (Barrett et al., 1997; Pérez-Barrales et al., 2006; Ferrero et al., 2009). Based on controlled hand pollinations, Goldblatt and Bernhardt (1990) have reported that pollen tubes could grow along the styles of either morph, independently of the morph from where the pollen was coming, even self-pollin. Nonetheless, it has been shown that some heterostylos and stigma-height dimorphic species present a late-acting type of incompatibility system (Dulberger, 1964; Phillip and Schou, 1981; Sage et al., 1999). Nivenia could have a similar system as indicated by the results of Ornduff (1983) on N. corymbosa, but this still needs to be ascertained in order to know if it stands for the observed isoplethy in the genus. Since the size of pollen grains is different between morphs (Goldblatt, 1993; present data), at least it was possible to explore if the observed polymorphism is functional in promoting disassortative pollen transfer, irrespective of the mating system in the populations. Unfortunately those differences are small enough to prevent the source (morph) of each individual pollen grain being unequivocally ascertained, as it was the case in a number of other studies (for some examples, see Ganders, 1979; Dulberger, 1992).
leading to isoplethic morph ratios. Moreover, there is some indication that the interplay between a restrictive floral morphology (deep and narrow tubes) and pollinator type and behaviour may promote this disassortative mating, as the data on flower phenotypic integration seem to point out. It is worth noting that N. parviflora (not included in this study) has a generalist pollinator array (Goldblatt and Manning, 2006) and much shorter flower tubes. On the other hand, N. fruticosa, even though monomorphic, has long corolla tubes and high floral integration values, which probably indicates that such a specialized pollination is a necessary but not a sufficient condition for heterostyly to evolve. Although there are still many open questions, it has become clear that Nivenia constitutes an excellent model system for further prospective studies on the evolution and function of different flower morphologies, including variable reciprocal herkogamy and floral shapes, as well as their interaction with pollinators.

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**LITERATURE CITED**


