Complex cytogeographical patterns reveal a dynamic tetraploid–octoploid contact zone

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Abstract. The distribution of cytotypes in mixed-ploidy species is crucial for evaluating ecological processes involved in the establishment and evolution of polyploid taxa. Here, we use flow cytometry and chromosome counts to explore cytoype diversity and distributions within a tetraploid–octoploid contact zone. We then use niche modelling and ploidy seed screening to assess the roles of niche differentiation among cytotypes and reproductive interactions, respectively, in promoting cytoype coexistence. Two cytotypes, tetraploids and octoploids, were dominant within the contact zone. They were most often distributed parapatrically or allopatrically, resulting in high geographic isolation. Still, 16.7% of localities comprised two or more cytotypes, including the intermediate hexaploid cytoype. Tetraploids and octoploids had high environmental niche overlap and associated with similar climatic environments, suggesting they have similar ecological requirements. Given the geographical separation and habitat similarity among cytotypes, mixed-ploidy populations may be transitional and subject to the forces of minority cytoype exclusion which lead to pure-ploidy populations. However, seed ploidy analysis suggests that strong reproductive barriers may enforce assortative mating which favours stable cytoype coexistence. High cytogenetic diversity detected in the field suggests that unreduced gamete formation and hybridization events seem frequent in the studied polyploid complex and might be involved with the recurrent polyploid formation, governing, as well, the gene flow between cytogetic entities.

Keywords: Contact zone; distribution patterns; Gladiolus communis; hexaploid; hybridization; niche modelling; niche overlapping; octoploid; tetraploid.

Introduction

Polyploidization, the duplication of complete chromosome sets, is widely considered an important mechanism of plant evolution (Soltis and Soltis 1999; Jiao et al. 2011) and sympatric speciation (Otto and Whitton 2000; Soltis et al. 2010). Based on recent molecular and fossil studies, polyploidy has been linked with radiations in species diversity throughout evolutionary history (Soltis et al. 2009) and associated with 15 % of...
speciation events in extant angiosperms (Wood et al. 2009). Consequently, polyploidy is pervasive in flowering plants. The standing incidence of polyploid species is estimated at 35% (Wood et al. 2009), with higher values being observed in specific geographic regions such as the Mediterranean basin (ranging between 37 and 47%; Marques et al. 2017) and the Arctic region (69 and 87%; Brochmann et al. 2004).

The geographic distribution of polyploids is useful for inferring mechanisms of polyploid evolution, coexistence and divergence. The spatial arrangement of cytotypes in situ is the result of several interacting processes operating in natural populations including formation and migration; ecological preferences, and competitive and dispersal abilities; and reproductive interactions, among others (Petit et al. 1999; Levin 2002; Lexer and van Loo 2006). Cytotype distributions can be characterized as sympatric, parapatric or allopatric depending on whether the different cytotypes grow intermixed, adjacent or disjunct, respectively (Petit et al. 1999; and illustrated in Fig. 2 of Mallet et al. 2009, which can be applied to polyploid complexes). Theoretical models predict that within zones of sympatry, mixed-ploidy populations are expected to be rare and evolutionarily unstable because frequency-dependent selection will drive the exclusion of the minority cytotype (Levin 1975; Rodriguez 1996; Husband and Schemske 2000). Still, numerous studies have documented mixed-ploidy populations (reviewed in Husband et al. 2013; and examples below). The presence of multiple cytotypes in the same population can reflect either a transitory stage, in which neopolyploids are recurrently formed, or a persistent stage such as when cytotypes are ecologically and reproductively isolated on a small spatial scale (e.g. Kolář et al. 2009; Jersáková et al. 2010). In this context, assessing the distribution of cytotypes within and among natural populations is crucial to build and test hypotheses that account for the successful establishment of polyploids.

Contact zones, areas with two or more cytotypes growing in close proximity, are thus considered natural laboratories within which to study evolutionary transitions through polyploidy. In recent years, an increasing number of studies have provided insights into ploidy-mediated processes occurring in contact zones (e.g. Husband et al. 2013; Ramsey and Ramsey 2014). Significant advances in this field have been largely fuelled by the ability to rapidly and easily screen thousands of individuals using flow cytometry (Kron et al. 2007). This approach has resulted in a proliferation of cytogeographical studies (e.g. Baack 2004; Kolář et al. 2009; Stählberg 2009; Trávníček et al. 2010; Castro et al. 2012; Zozomová-Lihová et al. 2015; Wefferling et al. 2017; reviewed in Ramsey and Ramsey 2014), which detect extensive cytogenetic diversity and, in several cases, occurrence of mixed-ploidy populations (e.g. Baack 2004; Kolář et al. 2009; Trávníček et al. 2010; Castro et al. 2012; Zozomová-Lihová et al. 2015; Wefferling et al. 2017), rare cytotypes (e.g. Kolář et al. 2009; Trávníček et al. 2010), production of unreduced gametes (e.g. Maceira et al. 1992; Burton and Husband 2001; Ramsey 2007; Castro et al. 2016a) or recurrent occurrence of gene flow (e.g. Husband 2004; Kolář et al. 2009; Castro et al. 2011). Particularly interesting are polyploid complexes with higher ploidies, such as diploid–hexaploid (e.g. Aster amellus; Castro et al. 2012) or tetraploid–octoploid complexes (e.g. Gymnadenia conopsea; Jersáková et al. 2010), that can produce even-ploidy hybrids, which are potentially more stable and lead to highly dynamic contact zones. Regardless of the increasing number of studies at contact zones, the available information is still scarce and insufficient for many plant groups and regions (Soltis et al. 2010, 2016; Marques et al. 2017).

Gladiolus communis is a Mediterranean polyploid complex with high morphological variation (Alonso and Crespo 2010). Multiple ploidy levels have been described for the complex, namely tetraploids (2n = 4x = 60 chromosomes; Fernandes et al. 1948; Fernandes 1950; Nilsson and Lassen 1971; Queirós 1980; Fernández et al. 1985) and octoploids (2n = 8x = 120 chromosomes; Fernandes and Queirós 1971; Løve and Kjellqvist 1973; Queirós 1980), although hexaploids (2n = 6x = 90) and duodecaploids (2n = 12x = 180) have also been occasionally reported in the Mediterranean basin (Darlington and Wylie 1955). The Iberian Peninsula seems to harbour this diversity (Fernandes et al. 1948; Fernandes and Queirós 1971; Queirós 1979) and areas of close contact between tetraploids and octoploids have been detected, for example, in calcareous regions from Central Portugal (Castro et al. 2016b). Occasionally, G. communis grows with another congeneric species, namely G. italicus, which, in the Iberian Peninsula, is represented by duodecaploid individuals (Queirós 1979; Pérez and Pastor Díaz 1994; although octoploids have also been described in the Mediterranean basin, e.g. Susnik and Lovka 1973; Strid and Franzen 1981; van Raamsdonk and De Vries 1989; Kamari et al. 2001). The high morphological variation of the group has led taxonomists to accept multiple taxonomic entities within the G. communis complex (e.g. Gussoni 1832; van Raamsdonk and De Vries 1989), although morphologically intermediate forms are found in natural populations, and many characters used to distinguish each taxon are extremely variable and largely overlap, even within populations (Hamilton 1980; revised in Alonso and Crespo 2010). Consequently, recent morphological reviews and preliminary molecular analyses failed to support the previous taxonomic delimitations and the species is currently accepted...
as a complex formed by three ploidy levels (Buchanan 2008; Alonso and Crespo 2010). Regardless of the variability detected in the species, nothing is known about the role of genome duplications generating diversity within this polyploid complex. Exploring cytotype diversity and distribution patterns, especially at contact zones, is thus crucial to understand ecological processes, such as ecological preferences and reproductive interactions, driving current diversity patterns at natural contact zones.

In this study, we explore in detail the cytotype diversity and distribution patterns in a tetraploid–octoploid Gladiolus communis contact zone. In particular, we pose the following specific questions: (i) what are the dominant cytotypes and how are they distributed in the contact zone? (ii) do cytotypes coexist and at which spatial scale? (iii) is coexistence facilitated by differences in environmental associations between cytotypes? And finally, (iv) is there evidence for the production of unreduced gametes and/or cytotype hybridization? To address our questions, cytotype diversity was studied at several spatial scales, namely (i) across the contact zone, to characterize the most dominant cytotypes and their environmental preferences within areas of contact; (ii) within mixed-ploidy populations, to measure microhabitat segregation; and (iii) among offspring from plants in pure- and mixed-ploidy populations, to detect cytotype diversity at early stages. Flow cytometric analyses complemented with chromosome counts were used to assess ploidy levels of all the sampled individuals. The reproductive success of pure- and mixed-ploidy populations was also quantified in natural conditions to depict fitness differences between cytotypes. The spatial arrangement of cytotypes in the contact zone was analysed with niche modelling tools to determine if differences in environmental requirements could explain cytotype distribution. If cytotypes differ in environmental requirements, we expect a mosaic contact zone with tetraploids and octoploids fairly isolated within a given spatial scale and with plants growing in different habitats or microhabitats. If no environmental differences are observed, we expect a tension zone where sympatric cytotype co-occurrence is possible, where intermediate cytotypes are detected and where other processes such as reproductive barriers, competition or dispersal abilities are expected to play major roles in driving distribution patterns.

Methods

Study system and studied region

Gladiolus communis is a perennial species that is widespread on the Iberian Peninsula and throughout the Mediterranean basin. The species produces an ovoid bulb, relatively thick roots, a cylindrical glabrous stem and linear leaves with typical parallel ribs. The pink bisexual flowers are zygomorphic and usually grouped in one spikelike inflorescence per individual. A second Gladiolus species, G. italicus, is found on the Iberian Peninsula and occurs in sympathy with G. communis in some places. Although very similar morphologically, these two species are easily distinguished based on inflorescence architecture, anther and filament lengths, and seed morphology. Gladiolus communis has a unilateral inflorescence, anthers equaling or shorter than the filaments, and broadly winged seeds, while G. italicus usually has a weakly distichous inflorescence, anthers longer than the filaments, and polyhedric apterous seeds (Hamilton 1980; Alonso and Crespo 2010).

In the Iberian Peninsula, G. communis is recognized as a polyploid complex comprising tetraploids (2n = 4x = 60 chromosomes), hexaploids (2n = 6x = 90) and octoploids (2n = 8x = 120) (e.g. Fernandes et al. 1948; Fernandes and Queirós 1971; Alonso and Crespo 2010) with duodecaploids being described elsewhere in the Mediterranean region (Darlington and Wylie 1955). The high morphological resemblance among G. communis cytotypes (Alonso and Crespo 2010; Cantor and Tolety 2011) suggests a putative autoploidy origin. The species is common in the calcareous regions from Central Portugal, where preliminary field sampling revealed the presence of tetraploid and octoploid populations growing in close proximity. This study focused on this contact zone, an area extending from 39.3° to 40.6° in latitude, and from 7.8° to 9.4° in longitude. This territory is dominated by calcareous rocks and presents a Mediterranean climate that exhibits a strong influence from the Atlantic Ocean, an attribute identified on the significant values of annual precipitation (1000–1300 mm). However, the dominance of poor soils determines a low water storage capacity, which, combined with a long and hot summer, determines the dominance of evergreen vegetation. Allied to such climatic conditions, human pressure contributed to current dominance of shrubby communities in the landscape, and constrained forests (evergreen and semi-deciduous) to very small patches, favouring the wide presence of open habitats. These open habitats are also characterized by the presence of limestone outcrops exposed to stressful ecological conditions that limit the installation of higher vegetation covers.

Although not exhaustive, additional sampling was extended beyond this area to determine the dominant cytotypes within the species. Also, because G. communis coexist with G. italicus, hybridization might occur and generate additional cytogenetic diversity, the duodecaploid G. italicus (Queirós 1979; Pérez and Pastor Díaz 1994) was also sampled whenever growing with G. communis.
Field sampling
Field collections were carried out during the flowering and fruiting seasons (mid-April to July) of *G. communis* from 2012 to 2015. Individual plants or clusters of plants were easily detected when blooming because of the tall inflorescences growing above the remaining vegetation. We sampled 81 populations across the contact zone where both tetraploid and octoploid cytotypes have been previously detected in close proximity. An additional group of 27 populations covering the western distribution of the species around the contact zone was also sampled to depict the dominant cytotypes [see Supporting Information—Table S1]. In each of the 108 populations, we collected ~3 cm² of fresh leaf of up to 53 individuals of *G. communis* (with an average of 20 individuals per locality, excluding two particularly large localities where more intensive sampling was done, with 106 and 454 plants being screened), and of *G. italicus* whenever detected growing with *G. communis* (up to 32 individuals, averaging 13 plants per locality). The sampled individuals were randomly selected, covering the extension of the population. Leaves were stored in labelled hermetic bags and maintained at 4 °C for later flow cytometric analysis (see section Genome size and DNA ploidy level estimates). Geographic coordinates of the population were recorded. Bulbs of nine localities identified in preliminary surveys as DNA tetraploid, DNA hexaploid and DNA octoploid populations (Castro et al. 2016b) were also collected, potted and maintained at the common garden for chromosome counts (see section Chromosome counts).

In addition, we sampled in mixed-ploidy populations more intensively to test for microhabitat segregation. Three mixed-ploidy populations containing tetraploids, hexaploids and/or octoploids were revisited and all adult, individuals (both vegetative and reproductive individuals) were mapped with *x/y* coordinates, tagged and sampled for ploidy level analyses using flow cytometry (see section Genome size and DNA ploidy level estimates). To delimit the clusters of plants growing in sympathy, screenings for *Gladiolus* plants were made around a radius of over 150 m around the cluster of plants initially detected or until an anthropogenic or natural barrier was observed. Additional mixed-ploidy populations were not sampled because they were disturbed by grazing or human activities.

Finally, we screened offspring from plants in pure- and mixed-ploidy populations to examine the production of unreduced gametes and/or hybridization events by the detection of rare cytotypes that might not reach the adult stage. For this, four tetraploid, two hexaploid, four octoploid and one mixed tetraploid–octoploid populations were revisited and individual plants with known ploidy were sampled to determine reproductive success and screen ploidy of the seeds (see section Reproductive success in natural populations).

Chromosome counts
Chromosome counts were used to calibrate genome size estimates, obtained using flow cytometry, to a given ploidy level. For this, the plants grown from bulbs collected in the selected natural populations and maintained in the common garden were used simultaneously for genome size estimates and chromosome counting. For chromosome counts, we followed the protocol of Goldblatt and Takei (1993), with some adjustments. Briefly, actively growing root tips were harvested and pretreated in 0.002 M aqueous 8-hydroquinoline at room temperature for 1630 h, and fixed in 95 % ethanol and glacial acetic acid (in a ratio of 3:1) for at least 48 h at 4 °C. Roots tips were hydrolysed in 1 M hydrogen chloride at 60 °C in a sand bath for 40 min, submerged in Schiff reagent (Greilhuber and Ebert 1994) for 1330 h, washed in sulphur water for three periods of 10 min and finally squashed under a glass cover in a drop of acetic orcein 2 %. Chromosome spreads were observed using a Nikon Eclipse 80i light microscope and photographed using a Nikon Plan Apo VC 100×/1.40 oil-immersion lens with a Q Imaging Retiga 2000R Fast 1394 digital camera and Q-Capture Pro v.7 software. A total of 40 individuals from nine populations were used to access chromosome number and genome size: 4x—populations MC147 (n = 10 individuals), MC193 (n = 1), MC195 (n = 4), MC201 (n = 1) and MC212 (n = 2); 6x—population MC211 (n = 4); 8x—populations MC032 (n = 8), MC143 (n = 3), MC190 (n = 4), MC193 (n = 2) and MC201 (n = 1) [see Supporting Information—Table S1].

Genome size and DNA ploidy level estimates
To estimate genome size and DNA ploidy levels, fresh leaves collected in natural populations were analysed using flow cytometry. Nuclear suspensions were prepared following Galbraith et al. (1983) by chopping the plant material of the sampled species together with leaf tissue of an internal reference standard. In the case of *Gladiolus* nuclear suspensions, 100 mg of leaf tissue or 2–5 seeds were co-chopped with 50 mg of leaf of *Solanum lycopersicum* ‘Stupické’ (2C = 1.96 pg; Doležel et al. 1992) or *Pisum sativum* ‘Ctirad’ (2C = 9.09 pg; Doležel et al. 1998). *Solanum lycopersicum* was used as the internal standard in most cases, except when unavailable, with *P. sativum* being used in those situations. Sample and standard were co-chopped in 1 mL of WPB buffer (WPB: 0.2 M Tris–HCl, 4 mM MgCl₂·6H₂O, 1 % Triton X-100, 2 mM EDTA Na₂·2H₂O, 86 mM NaCl, 10 mM metabolisfite, 1 % PVP-10, pH adjusted to 7.5 and stored at 4 °C;
using a razor blade. The resulting nuclear suspension was filtered through a 50 µm nylon filter and 50 µg ml⁻¹ propidium iodide (Fluka, Buchs, Switzerland) and 50 µg ml⁻¹ RNAse (Fluka) were added to the sample, to stain the DNA and avoid staining of double-stranded RNA, respectively. After 5 min of incubation, DNA fluorescence of the sample was analysed using a Partec CyFlow Space flow cytometer (S32 nm green solid-state laser, operating at 30 mW; Partec GmbH, Görlitz, Germany). Using Partec FloMax software v2.4d (Partec GmbH, Münster, Germany) the following four histograms were obtained: fluorescence pulse integral in linear scale (FL); forward light scatter (FS) vs. side light scatter (SS), both in logarithmic (log) scale; FL vs. time; and FL vs. SS in log scale [see Supporting Information—Fig. S1]. To digitally remove some of the debris, the FL histogram was gated using a polygonal region defined in the FL vs. SS histogram (see R1 in Supporting Information—Fig. S1), and was further applied to all the other graphics. At least 1300 nuclei in both sample and standard G₁ peaks were analysed per sample (Suda et al. 2007). Only coefficient of variation (CV) values of G₁ peak of G. communis below 5 % were considered acceptable (see examples in Supporting Information—Fig. S1), otherwise a new sample was prepared and analysed until this quality standard was achieved (Greilhuber et al. 2007).

Genome size was estimated in 41 populations by analysing three plants per population and cytotype individually (rarely less, unless there were no more plants in the locality, while in a few populations up to 30 individuals were analysed for genome size) [see Supporting Information—Table S2]. For the remaining individuals and populations, only DNA ploidy level information was gathered following the pooled sample strategy (5–6 individuals plus the reference standard). A total of 108 natural populations of G. communis and 2665 individuals were sampled and analysed [see Supporting Information—Table S1].

We used flow cytometry to measure DNA ploidy of offspring produced by plants of known ploidy. A total of 1252 seeds from 178 individuals from four tetraploid, two hexaploid and four octoploid pure-ploidy populations and one tetraploid–octoploid mixed population were analysed. We sampled 10–15 seeds per maternal individual, and 7–15 individuals per population and cytotype. For pure-ploidy populations of tetraploids and octoploids and mixed-ploidy population, five seeds were chopped simultaneously with the internal reference standard (pooled sample strategy) following the protocol described above, producing easy to interpret histograms. When analysing the seeds, at least two peaks (plus the peak of the internal standard) were always obtained, corresponding to the peak of the embryo and that of the endosperm. Consequently, the interpretation of each histogram was made with particular caution, determining the ploidy levels of all the peaks obtained in the histogram. Preliminary analyses revealed that hexaploid populations presented higher variability and thus only two seeds were pooled, in order to unambiguously assign the DNA ploidy levels of each seed.

The holoploid genome size (2C in pg; sensu Greilhuber et al. 2005) was obtained using the following formula:

\[
\text{Holoploid genome size (pg)} = \frac{G.\text{communis G₁ peak mean}}{\text{reference standard G₁ peak mean}} \times \text{standard genome size}
\]

Based on the chromosome counts obtained in this study and respective genome sizes, as well as the four chromosome numbers described in the literature for G. communis and G. italicus, DNA ploidy levels were inferred for each sample and individual. Populations were then characterized according to their DNA ploidy composition.

Descriptive statistics of holoploid genome size were calculated for each cytotype and species (mean, standard deviation of the mean, coefficient of variation of the mean, maximum and minimum values) based on the individual flow cytometric estimates. Mean and standard deviation of the mean were also calculated for the monoploid genome size (1C; holoploid genome size divided by inferred DNA ploidy level; sensu Greilhuber et al. 2005). Differences in holoploid and monoploid genome sizes among species and cytotypes were investigated using linear models (hereafter LM) performed in R software version 3.0.1 (R Core Development Team 2016), using the packages ‘car’ for Type-III analysis of variance (Fox et al. 2015), ‘lme4’ for generalized linear models (GLMs; Bates et al. 2014) and ‘multcomp’ for multiple comparisons after Type-III analysis of variance (Hothorn et al. 2017).

The geographical isolation index (GI) between the two dominant cytotypes (i.e. tetraploids and octoploids) at the contact zone was calculated according to the following formula (Husband et al. 2016), where only pure-ploidy and mixed-ploidy populations of tetraploids and octoploids from the contact zone were considered:

\[
\text{GI} = 1 - \frac{\text{no. mixed-ploidy populations}}{\text{total no. of populations}}
\]

**Environmental preferences**

The environmental associations of the two dominant cytotypes were evaluated through GLM, and spatial predictive models were produced based on niche modelling tools, aiming to assess niche overlap. To explore niche...
overlapping, two approaches were used considering two different spatial scales: (i) one with an extension encompassing the contact zone in Central Portugal; and (ii) the other extension encompassing the entire territory of mainland Portugal.

Variables were extracted from the following sources with a resolution of ~111 m: (i) bioclimatological data from http://home.isa.utl.pt/~tmh/aboutme/Informacao_bioclimatologica.html (methodology to obtain variables in Monteiro-Henriques et al. 2016); and (ii) data for soil conditions from: http://epic-webgis-portugal.isa.ulisboa.pt/. Values for climatic and soil variables were extracted for all the surveyed populations using the R package ‘dismo’ (Hijmans et al. 2017). Then, GLMs were used to explore climatic and soil variables and assess differences between tetraploid and octoploid populations (Table 1), namely for climatic variables [mean annual total precipitation (PP), mean temperature of the hottest month of the year (Tmax), mean temperature of the coldest month of the year (Tmin), mean maximum temperature of the coldest month of the year (M), mean minimum temperature of the coldest month of the year (m)], bioclimatic indexes [continentality index (IC), compensated thermicity index (ITC), summer ombrothermic index (Ios3)], soil conditions [texture (txt) and pH] and altitude. Correlation between variables was explored using Pearson coefficient for continuous variables and Spearman’s ρ for categorical variables, to assist variable selection by removing variables with correlation values higher than 0.7. The final set of variables selected included the following four which were also important descriptors of the type of habitat where the species grows: mean annual total precipitation, mean temperature of the hottest month, soil texture and pH (highlighted in bold in Table 1).

Spatial predictive models were calibrated based on presence/absence records collected in the field and the selected environmental and soil variables (Table 1). For the tetraploid data set, tetraploid populations were recorded as presences and octoploid populations as absences, and vice versa for the octoploid data set. Mixed tetraploid–octoploid populations were considered as presences for both cytotypes. For the contact zone (Central Portugal) we used data from 76 sampling points (including 33 tetraploid, 40 octoploid and 3 tetraploid–octoploid populations), corresponding to all the known occurrences of G. communis with a minimum distance of 600 m. For the territory of Portugal, and aiming to reduce the bias effect of spatial clustering associated with our intense screening in the contact zone, only occurrences that had a minimum distance of 10 km between them were selected, based

### Table 1. Characterization of the climatic and soil variables for tetraploid and octoploid populations of Gladiolus communis in the contact zone of Central Portugal. The mean, standard error of the mean (SE) and statistical tests (comparison between cytotypes) are provided for each variable and cytotype. Significance levels: ****P < 0.01; **0.05 < P < 0.01; n.s., non-significant. In bold the variables used in niche modelling are highlighted. Variables were extracted from the following sources with a resolution of ~111 m: (i) bioclimatological data from http://home.isa.utl.pt/~tmh/aboutme/Informacao_bioclimatologica.html (methodology to obtain variables in Monteiro-Henriques et al. 2016); and (ii) data for soil conditions from: http://epic-webgis-portugal.isa.ulisboa.pt/.

| Variables                      | Code | Tetraploid Mean ± SE, n = 43 | Octoploid Mean ± SE, n = 36 | ANOVA F, P value |
|-------------------------------|------|-----------------------------|----------------------------|-----------------|
| Precipitation PP              | 1096.11 ± 21.97a | 1106.89 ± 21.74a | 0.12 n.s. |
| Mean temperature of the hottest month of the year Tmax | 20.51 ± 0.13a | 20.95 ± 0.19b | 4.04* |
| Mean temperature of the coldest month of the year Tmin | 9.06 ± 0.10a | 8.91 ± 0.12a | 1.10 n.s. |
| Mean max. temp. of the coldest month of the year M | 13.52 ± 0.11a | 13.42 ± 0.12a | 0.38 n.s. |
| Mean min. temp. of the coldest month of the year m | 4.61 ± 0.08a | 4.50 ± 0.09a | 0.84 n.s. |
| Continentality index IC       | 11.44 ± 0.13a | 12.04 ± 0.19a | 6.95* |
| Compensated thermicity index ITC | 327.12 ± 2.78a | 325.81 ± 3.22a | 0.10 n.s. |
| Summer ombrothermic index Ios3 | 1.10 ± 0.03a | 1.09 ± 0.03a | 0.05 n.s. |
| Soil texture Texture          | 2.14 ± 0.29a | 2.25 ± 0.13a | 0.11 n.s. |
| Soil pH                       | 308.60 ± 46.51a | 99.44 ± 19.69b | 15.02*** |
| Altitude                      | 198.61 ± 18.63a | 169.23 ± 17.38a | 0.94 n.s. |
| Latitude                      | −8.47 ± 0.05a | −8.58 ± 0.03a | 3.14 n.s. |
| Longitude                     | 39.98 ± 0.05a | 40.01 ± 0.04a | 0.18 n.s. |

For the territory of Portugal, and aiming to reduce the bias effect of spatial clustering associated with our intense screening in the contact zone, only occurrences that had a minimum distance of 10 km between them were selected, based
on radon selection, resulting in a subset of 66 sampling points (including 35 tetraploid, 19 octoploid and 6 tetraploid–octoploid populations).

Environmental niche modelling (ENM) of tetraploids and octoploids was created using R package ‘biomod2’ (Thuiller et al. 2016). Final model for each cytotype is based on the combination of results from different modelling techniques, each one replicated 30 times after data splitting into training (70 %) and testing (30 %) subsets based on random selection, aiming to reduce uncertainty and to produce robust models (Phillips et al. 2006; Araújo and New 2007). In the resampling replication, each specific occurrence was used only once in each run, as training or as test without replacement, making all replicates statistically independent (Phillips 2008). Models were evaluated based on the independent accuracy measure AUC of ROC (area under the curve of the receiver operating characteristic), and only those with AUC > 0.7 were used in the ensemble forecasting procedure, the approach used to produce the final model for each cytotype.

Model evaluation revealed high ROC values (contact zone: 4x—0.79 ± 0.01 and 8x—0.79 ± 0.01; Portugal: 4x—0.77 ± 0.01 and 8x—0.76 ± 0.01) and relatively low omission rates (contact zone: 4x—0.19 ± 0.02 and 8x—0.28 ± 0.02; Portugal: 4x—0.23 ± 0.01 and 8x—0.28 ± 0.01). However, when considering the binary projections, the omission rates decrease to 0.10 and 0.09 for the tetraploid and octoploid models in the contact zone, respectively, and 0.17 and 0.04 in Portugal (tetraploids and octoploids, respectively), demonstrating that the models were able to predict the occurrences with high accuracy, namely for octoploids. The binary projection produced by the final model of each cytotype was used to calculate niche overlap.

Cytotype niche overlap was quantified through the metric of proportional similarity of the distribution of both cytotypes, using Schoener’s D (a measure of niche similarity; Schoener 1970). This metric ranges from zero (no overlap) to one (complete overlap). The ‘ecospat’ (Broennimann et al. 2012) and ‘raster’ (Hijmans et al. 2017) packages were used to perform niche identity and similarity tests (Warren et al. 2008; Broennimann et al. 2012). In niche equivalency (identity test), the points of both cytotypes were pooled and randomly split in two groups according to size of the original data set. This new data set was used in D calculation, and the process was repeated 100 times (to obtain confidence intervals that enable evaluation of the null hypothesis). The resulting D values (simulated values) were compared with the observed D value, and cytotype niches were considered equivalent if the observed D value fell within the 95th percentile of the simulated D values (Broennimann et al. 2012). In niche similarity (similarity test), we evaluate if the environmental niches of the two cytotypes were distinguishable from each other. In this case, the comparison was between the points of one cytotype and random points from the geographic range of the other cytotype. As in the identity test, the process was repeated 100 times and D values were calculated. The results revealed if niche overlap between the cytotypes is greater (niche conservation) or lower (niche divergence) than expected, according to the geographic region of the other cytotype. All the models and analyses were performed in R software version 3.0.1 (R Core Development Team 2016).

Reproductive success in natural populations

The reproductive success of each cytotype was evaluated in 11 natural populations, namely 10 pure-ploidy populations (including four tetraploid, two hexaploid and four octoploid populations) and one mixed-ploidy population composed by tetraploid and octoploid individuals (MC201). In each population, 11–20 individuals of known ploidy level were labelled and infructescences collected in individually labelled bags. The number of fruits was counted for each inflorescence and fruit set calculated as the proportion of flowers that developed into fruit. The number of morphologically viable seeds (based on their size and shape) was assessed in all fruits, and the seed:ovule ratio (S:O ratio) was calculated by dividing the number of morphologically viable seeds by the number of ovules. The total reproductive success of populations and cytotypes was also calculated by multiplying the S:O ratio by the fruit set. Descriptive statistics were calculated for each population type.

Differences in fruit set, S:O ratio and total reproductive success between the three cytotypes (tetraploids, hexaploids and octoploids) within pure-ploidy populations, differences between tetraploids and octoploids in the mixed-ploidy population, and differences between pure- and mixed-ploidy populations (excluding hexaploid ones) were assessed using GLM. Mixed models with individual and population as random factors were initially used, but the random factors were further removed due to low variance in comparison with residuals (Bolker et al. 2009). A binomial distribution with a logit link function was used for fruit set, and a Gaussian distribution with an identity link function was used for S:O ratio and total reproductive success after transformation with the arcsine of the square root. When significant differences were obtained, post hoc tests for multiple comparisons were performed.

All analyses were performed in R software version 3.0.1 (R Core Development Team 2016), using the packages ‘car’ for Type-III analysis of variance...
Castro et al. – Cytogeographical patterns in a tetraploid–octoploid complex

(Fox and Weisberg 2015), ‘lme4’ for GLMs (Bates et al. 2014) and ‘multcomp’ for multiple comparisons after Type-III analysis of variance (Hothorn et al. 2017).

Results

Genome size and cytogenetic diversity

Based on chromosome counts and flow cytometric analyses, we detected three ploidy levels in G. communis: tetraploids with $2n = 4x = 60$ chromosomes (Fig. 1A) and an average genome size of $2.69 \pm 0.06$ pg/2C (mean ± SD); hexaploids with $2n = 6x = 90$ chromosomes (Fig. 1B) and an average genome size of $4.07 \pm 0.07$ pg/2C; and octoploids with $2n = 8x = 120$ chromosomes (Fig. 1C) and an average genome size of $5.42 \pm 0.14$ pg/2C (Table 2; [see Supporting Information—Table S2]).

Genome size estimates also suggest the occurrence of nonaploid G. communis individuals, characterized by genome sizes with nine times the monoploid genome size ($1C$) values obtained for the other ploidy levels, and had mean genome size of $6.10 \pm 0.18$ pg/2C (Table 2; Fig. 2C). These individuals were rare and we were unable to confirm their ploidy using chromosome counts. Gladiolus italicus had a higher genome size (2C = $7.27 \pm 0.17$ pg) than G. communis, consistent with duodecaploids, as described for the species (Table 2; Fig. 2D). The holoploid genome sizes (2C) of the five cytotypes differed significantly ($F_{4, 175} = 7691.3, P < 0.001$; Table 2). Monoploid genome size values were conserved within G. communis, with no significant differences being observed between cytotypes ($F_{3, 155} = 7691.3, P = 0.5272$; Table 2). However, monoploid genome size of G. communis (0.67 ± 0.03 pg) was significantly higher than for G. italicus (0.61 ± 0.01 pg; $F_{1, 178} = 7691.3, P < 0.001$).

Geographic distribution of cytotypes

Tetraploids and octoploids were prevalent across the geographic area sampled, both occurring in pure- and in mixed-ploidy populations (Fig. 3). No marked segregation pattern of cytotype arrangement in space was observed: tetraploids seem to occur across the entire area surveyed, and octoploids in the centre and south of the surveyed area, forming broad contact zones. Minority cytotypes were also detected, namely hexaploids, which were observed growing with other cytotypes and occasionally found forming pure populations (Fig. 3). A few nonaploids in a mixed-ploidy population harbouring all cytotypes of G. communis were also detected (Fig. 3) [see Supporting Information—Table S3]. Most populations were cytogenetically uniform (i.e. pure-ploidy populations, 86.1 %) and, in the majority of cases, were composed of either tetraploid or octoploid individuals (43.5 and 39.8 %, respectively). Hexaploids were detected growing alone in three locations (2.8 %) (Fig. 3). Populations harbouring two or more cytotypes (i.e. mixed-ploidy populations) represented 13.9 % of all sampled populations. The mixed-ploidy populations presented different cytotype compositions: tetraploids and hexaploids (4.6 %), in which the former is more frequent than the latter; tetraploids and octoploids (5.6 %) again, in which tetraploids are generally more abundant than octoploids, except in one population; tetraploids, hexaploids, octoploids and nonaploids (0.9 %; one population), where octoploids are the dominant cytotype; and hexaploids and octoploids (2.8 %), in which octoploids are dominant, except in one location where only two plants, one of each cytotype, were found [see Supporting Information—Table S3].

Within the contact area (Fig. 3B), most localities contained a single ploidy of either tetraploids (42.0 %), octoploids (44.4 %) or rarely hexaploids (2.5 %). These populations were distributed mostly in parapatry; still, cytotypes were found growing in sympatry in some locations (11.1 %) (Fig. 3B). Octoploid populations occur from north to south, resulting in cytogenetically diverse contact zones with tetraploids to the east, south and southwest. At these contact zones, areas with different types of mixed-ploidy populations were detected. Hexaploids were frequent in the contact zones between tetraploids

Figure 1. Gladiolus communis chromosome counts. (A) tetraploid ($2n = 4x = 60$ chromosomes), (B) hexaploid ($2n = 6x = 90$) and (C) octoploid ($2n = 8x = 120$) individuals. Bar = 20 μm.
and octoploids, although they were also detected in other places of the screened area, growing with tetraploid individuals. Tetraploids and octoploids, the two main cytotypes, were observed growing together in four locations out of the 81 populations at the contact zone (4.9 %), resulting in a total GI of 0.95, with tetraploids and octoploids presenting a similar individual geographical isolation index (GI<sub>4x</sub> = 0.90, GI<sub>8x</sub> = 0.91).

The detailed screening of three selected mixed-ploidy populations revealed variable patterns of cytotype distribution within each population (Fig. 4). In the tetraploid–octoploid population (MC201), cytotypes were distributed in two well-defined clusters separated by >20 m, with tetraploids being restricted to the north-east side and octoploids to the south-west of the population (Fig. 4A). The mixed-ploidy population with tetraploids and hexaploids (MC232) was dominated by tetraploid individuals, with a few hexaploids growing intermingled (Fig. 4B). The population with the highest cytogenetic diversity (MC193) revealed to be dominated by octoploid individuals with a few tetraploid, hexaploid and nonaploid plants growing intermingled (Fig. 4C). While MC201 and MC193 were located in the contact zones, MC232 is located in an otherwise tetraploid zone (Fig. 3).

### Environmental preferences

Niche geographic overlap between tetraploids and octoploids at both the contact zone (Schoener’s D metric, D = 0.03) and Portugal (D = 0.01) was low (Table 3). However, and despite little geographical overlap, there was no statistical evidence that the environmental niches differed, i.e. neither niche equivalency nor niche similarity was rejected (Table 3). This indicates that environmental niche of the dominant cytotypes was equivalent within the suitable ranges of both tetraploids and octoploids, and that environmental niche of each cytotype was similar to the suitable range of the other cytotype. At the contact zone, the selected climatic and

### Table 2. Genome size and DNA ploidy level estimations for Gladiolus communis and G. italicus. Holoploid genome size (G.s.; 2C) is provided for each species and cytotype as mean and standard deviation (SD) in picograms (pg), followed by coefficient of variation (CV, %), DNA range (minimum, Min, and maximum, Max, genome size values); mean and standard deviation of the mean is also provided for the monoploid genome size (1Cx); total number of individuals (N total) and populations (N pop) analysed for genome size are also provided for each ploidy level. Chromosome numbers (Chr. number) for each species based on chromosome counts of this study and bibliographic records are also provided. DNA ploidy levels: tetraploid (4x), hexaploid (6x), octoploid (8x), nonaploid (9x) and duodecaploid (12x). Different letters denote statistically significant differences at P < 0.05. 1 Chromosome numbers detected in this study; 2 Chromosome counts documented in the bibliography; 3 DNA ploidy level extrapolated based on the genome size values obtained here and on the chromosome counts available from other ploidy levels.

| Species          | DNA ploidy level | Chr. number | Holoploid G.s. (2C, pg) | Monoploid G.s. (1Cx, pg) | N total | N pop |
|------------------|------------------|-------------|-------------------------|--------------------------|---------|-------|
|                  |                  |             | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max | Mean            | SD | CV (%) | Min | Max |
| G. communis      | 4x               | 60<sup>1,2</sup> | 2.69<sup>a</sup> | 0.06 | 2.28 | 2.58 | 2.86 | 0.67<sup>a</sup> | 0.02 | 57   | 16  |
| G. communis      | 6x               | 90<sup>1,2</sup> | 4.07<sup>b</sup> | 0.07 | 1.76 | 3.93 | 4.19 | 0.68<sup>b</sup> | 0.01 | 9    | 3   |
| G. communis      | 8x               | 120<sup>1,2</sup> | 5.42<sup>c</sup> | 0.14 | 2.53 | 5.13 | 5.73 | 0.68<sup>c</sup> | 0.02 | 91   | 21  |
| G. communis      | 9x               | 135<sup>3</sup> | 6.10<sup>d</sup> | 0.18 | 2.89 | 5.98 | 6.23 | 0.68<sup>d</sup> | 0.02 | 2    | 1   |
| G. italicus      | 12x              | 180<sup>2</sup> | 7.27<sup>e</sup> | 0.17 | 2.31 | 6.97 | 7.55 | 0.61<sup>e</sup> | 0.01 | 21   | 10  |

**Figure 2.** Flow cytometric histograms of relative propidium iodide fluorescence from nuclei isolated from fresh leaves of Solanum lycopersicum ‘Stupické’ (S.l.) and different cytotypes and/or species of Gladiolus: (A) tetraploid (4x) and hexaploid (6x), (B) octoploid (8x) and (C) nonaploid (9x) individuals of G. communis, and (D) duodecaploid (12x) individual of G. italicus.
soil variables explained 62.98% of the variance in the distribution (Fig. 5A), and a high environmental overlap of a given cytotype within the niche of the opposite cytotype was observed (74.87 and 61.95% for tetraploids and octoploids, respectively; Fig. 5B). A similar pattern was observed for Portugal, although the climatic and soil variables explained higher variance than at the contact zone (74.78%; Fig. 5C). A high environmental overlap between cytotypes was also observed (91.51 and 47.96% for tetraploids and octoploids, respectively; Fig. 5D).

Reproductive success in natural populations and offspring cytogenetic composition

Plants in all the natural populations successfully formed fruits and seeds. However, the success differed according to the cytotype and population type. Pure-ploidy populations (excluding the hexaploid populations) had higher reproductive success compared to the mixed-ploidy population for all parameters (fruit set: $F_{1,1033} = 15.51, P < 0.001$; S:O ratio: $F_{1,796} = 4.62, P = 0.032$; reproductive success: $F_{1,1033} = 21.04, P < 0.001$; Fig. 6). Within pure-ploidy populations, significant differences between cytotypes were observed for all the variables (fruit set: $F_{2,1087} = 4.96, P = 0.007$; S:O ratio: $F_{2,770} = 100.18, P < 0.001$; reproductive success: $F_{2,1087} = 28.34, P < 0.001$), with octoploids having lower fruit set than tetraploids. S:O ratio and reproductive success were similar in tetraploids and octoploids ($P > 0.05$), but significantly higher than for hexaploids ($P < 0.05$; Fig. 6B and C). Within the mixed-ploidy population, no significant differences were observed between the cytotypes for any of the reproductive variables (fruit set: $F_{1,79} = 0.27, P = 0.603$; S:O ratio: $F_{1,37} = 0.01, P = 0.934$; reproductive success: $F_{1,79} = 0.15, P = 0.698$).

The analyses of offspring ploidy revealed that tetraploid and octoploid individuals, in both pure-ploidy and mixed-ploidy populations, produced seeds with the same ploidy as the mother plants (Fig. 6C). Tetraploid plants in pure populations produced a few aneuploids (<1% of the offspring; Fig. 6C) [see Supporting Information—Table S4]. In contrast, the flow cytometric analyses of the seeds from hexaploid individuals pointed out highly variable genome sizes, the analyses of the genome size estimates suggest the following DNA ploidy levels: 62% of seeds were aneuploid, 20% were pentaploids and 18% were hexaploid, although further confirmation is needed.

Discussion

This study corroborates the existence of high cytogenetic diversity within the *G. communis* polyploid complex. Two
dominant cytotypes, tetraploids and octoploids, were observed along with two minority cytotypes, mostly hexaploids, and rarely nonaploids. Tetraploids and octoploids have been well documented on the Iberian Peninsula through chromosome counts (Fernandes et al. 1948; Fernandes 1950; Fernandes and Queirós 1971; Nilsson and Lassen 1971; Löve and Kjellqvist 1973; Queirós 1980; Fernández et al. 1985). Also, hexaploids have been previously reported in the Mediterranean basin (Darlington and Wylie 1955). We observed them in 11% of the sampled localities (12 of 105 localities), commonly growing with one of the dominant cytotypes and occasionally in pure-ploidy populations. Nonaploids are reported here for the first time and were detected in the most diverse mixed-ploidy population.

Despite the cytogenetic diversity reported in G. communis, almost nothing was known about the geographic distribution of the cytotypes, or the presence and structure of its contact zones. Based on our survey, tetraploids occurred throughout the sampling area, although they were more common in the north and central regions of Portugal. Octoploids occurred in south and central regions of Portugal, but not in the north, notwithstanding the fact that more extensive surveys are needed to confirm this pattern. Although several mixed-ploidy populations were found, the geographical isolation index between tetraploids and octoploids is high, reflecting the fact that most of the populations contain a single cytotype. These populations distribute in space allopatrically or parapatrically, forming several contact zones between tetraploids and octoploids. However, despite that tetraploids and octoploids have non-overlapping distributions, they can inhabit similar environmental niches. Niche identity and similarity tests showed that tetraploids and octoploids occupy similar niches and are not differentiated in their environmental niches, showing niche conservation. These results contrast with other polyploid complexes for which niche differentiation, driven either by the direct effects of polyploidy or by subsequent selection, underlies the spatial separation of cytotypes and allows them to escape the minority cytotype disadvantage (e.g. Glennon et al. 2012; Thompson et al. 2014; Visger et al. 2016; Muñoz-Pajares et al. 2017). Still, the absence of environmental niche differences might not be completely unexpected as polyploids might not differ from their lower ploidy ancestors, either because they have been formed recently and the new polyploids did not have time to diverge from their progenitors, because genome duplications did not generate

Table 3. Niche analyses in Gladiolus communis. For each region studied, equivalency (D and P values) and similarity (P value) tests for suitable habitat are given.

| Suitable habitat | Equivalence test | Similarity test (P values) |
|------------------|------------------|-----------------------------|
|                  | D value  | P value | Tetra -> Octo | Octo -> Tetra |
| Contact zone     | 0.034     | 0.960   | 0.406          | 0.337          |
| Portugal         | 0.009     | 0.515   | 0.535          | 0.515          |

Figure 4. Fine-scale distribution of Gladiolus communis individuals within three mixed-ploidy populations: (A) tetraploid and octoploid mixed-ploidy population (MC201), (B) tetraploid and hexaploid mixed-ploidy population (MC232); and (C) tetraploid, hexaploid, octoploid and nonaploid mixed-ploidy population (MC193). Each point represents one individual plant mapped in a x/y system where distance is given in metres (m): tetraploids (4x), hexaploids (6x), octoploids (8x) and nonaploids (9x) individuals are represent by white, grey, black and dark grey points, respectively.
significant direct physiological changes, and/or because they might have been subjected to recurrent gene flow (Godsoe et al. 2013; Laport et al. 2016). Also, the effect of other environmental parameters on the distribution patterns observed in *G. communis* cannot be completely ruled out, nor the fact that niche differentiation might occur at a special resolution higher than that used in our study, although we did not find any clear evidence of differentiation in the field, namely considering the type of vegetation or the type of substrate in the mixed-ploidy populations detected (M. Castro, field observations).

Considering that *G. communis* cytotypes do not differ in suitable habitat, there should be historical processes and other ecological determinants shaping their distributional patterns, similarly to what has been observed in several polyploid complexes (e.g. Baack 2004, 2005; Pannell et al. 2004; Baack and Stanton 2005; Godsoe et al. 2013; Münzbergová et al. 2013; Wefferling et al. 2017). Contact zones are generated by direct emergence of neopolyploids in lower ploidy parental populations or through secondary contact of previously allopatric distributions in which cytotypes colonized the area separately in dissimilar ways and at different timings (Petit et al. 1999; Lexer and van Loo 2006). Although we still do not know the origin of *G. communis* contact zones, the different cytotype compositions found in natural populations provide significant insights into the processes that might be occurring at these areas (e.g. Husband and Schemske 1998; reviewed in Husband et al. 2013; Suda et al. 2013). One of the main
observations is the fairly few mixed-ploidy populations (10 vs. 90 % of mixed- and pure-ploidy populations), all composed of unbalanced number of tetraploid and octoploid plants (either dominated by tetraploid or by octoploids). In the absence of environmental differences, and regardless of the origin of the contact zone, *G. communis* mixed tetraploid–octoploid populations are expected to be more common at contact areas than detected here (4.9 % in the contact zone and 6.5 % from the total), since cytotypes might disperse to areas of the other cytotype and/or new cytotypes might be formed. Consequently, the high geographical isolation observed between *G. communis* cytotypes suggests that the mixed-ploidy populations might be transitory because strong frequency-dependent selection is expected to eliminate the minority cytotype as a result of fitness disadvantage generated by its lower number. This selection will ultimately drive the occurrence of pure-ploidy populations at contact zones (*Levin 1975; Husband 2000*).

However, tetraploid–octoploid populations may persist in nature. The regular production of unreduced gametes and the presence of reproductive barriers promoting assortative mating might lessen the magnitude

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**Figure 6.** Reproductive fitness of natural pure- and mixed-ploidy populations of *Gladiolus communis*: (A) fruit set; (B) S:O ratio (number of viable seeds divided by the number of ovules); and (C) reproductive success (fruit set multiplied by S:O ratio). In (C) the proportion of DNA ploidy levels detected in the offspring is also given. DNA ploidy levels: tetraploid (4x), pentaploid (5x), hexaploid (6x), octoploid (8x); seeds with genome size values out of the range of variation of each ploidy levels were assumed as aneuploids (An.). Different letters correspond to statistically significant differences as follows: (i) differences between population type (pure- vs. mixed-ploidy populations, excluding 6x) are denoted by upper case letters; and (ii) differences between ploidy levels within population type (among 4x, 6x and 8x from pure populations, and between 4x and 8x from the mixed-ploidy population) are denoted by lower case letters (Tukey HSD; *P* < 0.05); n.s. correspond to non-significant differences (*P* > 0.05).
of frequency-dependent selection and enable cytotype coexistence (e.g. Felber 1991; Segraves and Thompson 1999; Husband 2004; Husband and Sabara 2004; Kennedy et al. 2006). Octoploids might emerge directly in tetraploid populations through the union of two unreduced gametes \((n = 4x)\) or might result from seed dispersal from neighbouring octoploid populations. Unreduced gamete production has been detected in controlled pollinations in tetraploid G. communis (M. Castro et al., in preparation) and in screenings in natural populations through the detection of hexaploid individuals (see below). The rates at which unreduced gametes are produced might feed the population of octaploids enabling their maintenance within tetraploid populations (Felber 1991; Husband 2004). Additionally, seed ploidy analyses in a tetraploid–octoploid population suggest that strong reproductive barriers may enforce assortative mating, further favouring cytotype coexistence. Reproductive barriers driven, for example, by phenological and/or morphological mismatch, different pollinator assemblages or preferences, and/or gametic isolation will, thus, play a major role for overcoming minority cytotype exclusion in mixed-ploidy populations. Therefore, the fate of octoploids might depend not only on the rates of unreduced gamete formation but also on the reproductive isolation levels. Additionally, differences in other traits, such as perenniality or asexual reproduction, could compensate for the minority cytotype disadvantage (e.g. Rodriguez 1996; Kao 2007; Castro et al. 2016a). In other polyploid complexes, traits such as the production of bulbs represented an advantage, enabling new cytotypes to persist at initial stages and spread within lower ploidy populations (e.g. Allium oleraceum; Duchoslav et al. 2010; G. × sulistrovicus; Szczepaniak et al. 2016). If, through some of these traits, the number of octoploids can surpass the number of tetraploids, at some time octoploids might even outcompete tetraploids and exclude them from the population, as observed in other polyploid complexes (e.g. Buggs and Pannell 2007). Indeed, octoploids were observed as the dominant cytotype in some mixed-ploidy populations of the contact zone. Future studies on the contribution of all the above-mentioned processes, and on the relative contribution of sexual vs. asexual reproduction for the maintenance of the populations, are needed to fully understand the dynamics of mixed-ploidy populations.

The cytotype composition of G. communis natural populations also revealed that hexaploid plants might be more common than previously thought. These hexaploids might have originated through two different pathways. Hexaploids may originate from tetraploids through the union of reduced \((n = 2x)\) and unreduced \((n = 4x)\) gametes (Ramsey and Schemske 1998). Indeed, unreduced gamete formation is an important pathway for new polyploid emergence and has been shown to be common in nature (Felber 1991; Bretagnolle and Thompson 1995; Ramsey and Schemske 1998; Husband 2004; Ramsey 2007). This might explain the detection of hexaploid plants frequently found in otherwise tetraploid populations. Alternatively, hexaploids may form as a result of hybridization events between tetraploid and octoploid G. communis individuals. Gladiolus communis is pollinated by generalist pollinators that seem to have no cytotype preferences and might move pollen within mixed-ploidy populations or between populations in close proximity (M. Castro et al., in preparation). Additionally, controlled pollinations between tetraploid and octoploid plants were also successful in producing hexaploid offspring (M. Castro et al., in preparation). Either one of these pathways, i.e. unreduced gamete formation or hybridization, may operate in natural populations, being difficult to distinguish them without genetic markers. However, the relative abundance of tetraploid–hexaploid populations and paucity of tetraploid–hexaploid–octoploid populations suggests that the majority of the hexaploids are formed through unreduced gametes in tetraploid populations. Additionally, unreduced gamete production has been frequently detected in controlled pollinations involving tetraploid G. communis (M. Castro et al., in preparation), supporting it as a probable pathway for new cytotype emergence in natural populations. Quantifying unreduced gamete production in natural populations will provide significant insights on how frequent this process could be involved with hexaploid emergence.

Interestingly, hexaploid individuals were also found forming pure-ploidy populations, showing that this cytotype can successfully establish and spread beyond parental populations, although their sexual reproductive fitness was revealed to be lower in comparison with tetraploids and octoploids. Regardless of the lower fitness, recurrent unreduced gamete formation and asexual reproduction might enable to compensate for this disadvantage (e.g. Husband 2004; Kao 2007; Castro et al. 2016a). The successful establishment of hexaploid plants further contributes to the diversification of the complex. Ultimately, contact zones result from the combination of several factors, including historical factors, unreduced gamete formation, pollen flow and hybridization events, and seed dispersal, among others (Petit et al. 1999; Levin 2002; Lexer and van Loo 2006). Future studies reconstructing the history of the complex and quantifying unreduced gamete production, and its ability to hybridize, would provide significant insights on the dynamics of the distribution of G. communis.
The genome size of *G. italicus* suggests that this species is duodecaploid in the studied area, which is in accordance with chromosome counts for the Iberian Peninsula (Queirós 1979; Pérez and Pastor Díaz 1994), and contrasts with the dominance of the octoploids elsewhere in the Mediterranean basin (Susnik and Lovka 1973; Strid and Franzen 1981; van Raamsdonk and de Vries 1989; Kamari et al. 2001). Interestingly, the variation in monoploid genome size within *G. communis* cytotypes was very low and differed significantly from that of *G. italicus* (~9%). Given the magnitude of the differences between *G. italicus* and *G. communis*, both in ploidy levels and in monoploid genome sizes, holoploid genome size might be an important tool to detect hybridization (e.g. Kolář et al. 2009; Agudo-García 2017). In our study, *G. italicus* and *G. communis* were found growing in sympatry in 13% of localities; however, all the *G. italicus* individuals were duodecaploid. In most of the cases, the duodecaploid *G. italicus* was found growing with the octoploid *G. communis* (12 out of 14 localities); still, no decaploids were observed in these localities. When growing with the tetraploid *G. communis*, no octoploid individuals with lower genome size resulting from the hybridization between the two species (~5.00 pg based on the monoploid genome sizes of each species) were observed. Although hybridization has been suggested to occur in these and in other *Gladiolus* species (e.g. van Raamsdonk and de Vries 1989; Mifsud and Hamilton 2013; Szczepaniak et al. 2016), we were not able to detect hybrids between *G. italicus* and *G. communis*. This suggests that, in the studied range, hybridization between them might be less common, either because of assortative mating or hybrid offspring inviability. Monoploid genome size also suggests a close relationship between the cytotypes of *G. communis*, pointing to an autoployploid origin of the complex in the studied area. This is also supported by the high morphological resemblance between *G. communis* cytotypes (Alonso and Crespo 2010; Cantor and Tolety 2011) and by the lack of evidence supporting hybridization between *G. communis* and *G. italicus* in this region. Still, the origin of *G. communis* polyploid complex needs to be properly evaluated in future studies.

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## Contributions by the Authors

S.C., B.H. and J.L. designed the research experiment; M.C., S.C. and J.L. developed the field and laboratory work; M.C. performed the statistical analyses; and M.C. and A.F. performed the niche modelling analyses; all the authors were involved in the discussion of the results and writing of the manuscript, approving the final document.

## Conflict of Interest

None declared.

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Conclusions

In this study, we find a complex cytogeographical pattern in *G. communis*, which opens several hypotheses that might explain the formation and maintenance of its tetraploid–octoploid contact zone. According to our results, tetraploids and octoploids do not differ in their environmental requirements, potentially growing in similar habitats. Without differences in habitat requirements, mixed-ploidy populations were expected to be frequent; however, a high geographical isolation index was obtained. The high geographical isolation observed in nature, along with habitat similarity, suggests that the cytotype distribution in *G. communis* reflects historical patterns of migration and colonization, and further selection against minority cytotype, and does not result from different environmental requirements, creating a tension zone of contact. Still, in areas of contact, reproductive barriers might mediate assortative mating and enable cytotype coexistence. Nevertheless, the high cytogenetic diversity detected in the field suggests that unreduced gamete formation and hybridization events seem frequent in this complex and might be involved with recurrent polyploid formation and with gene flow between cytogenetic entities. Future studies involving reciprocal transplants will provide significant insights into the dynamics of this polyploid complex.
Supporting Information
The following additional information is available in the online version of this article—

Figure S1. Flow cytometry graphics analysed for each sample.

Table S1. Geographic information of sampled Gladiolus populations.

Table S2. Genome size variation in Gladiolus communis.

Table S3. Mixed-ploidy populations of Gladiolus communis.

Table S4. DNA ploidy levels of the offspring of pure- and mixed-ploidy populations of Gladiolus communis.

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