Multiple late-Pleistocene colonisation events of the Antarctic pearlwort *Colobanthus quitensis* (Caryophyllaceae) reveal the recent arrival of native Antarctic vascular flora

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Abstract

**Aim:** Antarctica's remote and extreme terrestrial environments are inhabited by only two species of native vascular plants. We assessed genetic connectivity amongst Antarctic and South American populations of one of these species, *Colobanthus quitensis*, to determine its origin and age in Antarctica.

**Location:** Maritime Antarctic, sub-Antarctic islands, South America.  
**Taxon:** Antarctic pearlwort *Colobanthus quitensis* (Caryophyllaceae).

**Methods:** Four chloroplast markers and one nuclear marker were sequenced from 270 samples from a latitudinal transect spanning 21–68° S. Phylogeographic, population genetic and molecular dating analyses were used to assess the demographic history of *C. quitensis* and the age of the species in Antarctica.

**Results:** Maritime Antarctic populations consisted of two different haplotype clusters, occupying the northern and southern Maritime Antarctic. Molecular dating analyses suggested *C. quitensis* to be a young (<1 Ma) species, with contemporary population structure derived since the late-Pleistocene.

**Main conclusions:** The Maritime Antarctic populations likely derived from two independent, late-Pleistocene dispersal events. Both clusters shared haplotypes with sub-Antarctic South Georgia, suggesting higher connectivity across the Southern Ocean than previously thought. The overall findings of multiple colonization events by a vascular plant species to Antarctica, and the recent timing of these events, are of significance with respect to future colonizations of the Antarctic Peninsula by vascular plants, particularly with predicted increases in ice-free land in this area. This study fills a significant gap in our knowledge of the age of the contemporary Antarctic...
terrestrial biota. Adding to previous inferences on the other Antarctic vascular plant species (the grass Deschampsia antarctica), we suggest that both angiosperm species are likely to have arrived on a recent (late-Pleistocene) time-scale. While most major groups of Antarctic terrestrial biota include examples of much longer-term Antarctic persistence, the vascular flora stands out as the first identified terrestrial group that appears to be of recent origin.

KEYWORDS
angiosperm, Antarctica, biogeography, dispersal, island, pearlwort, South America, Southern Ocean

1 | INTRODUCTION

Antarctic terrestrial ecosystems experience some of the most extreme conditions on Earth. Estimates of current ice-free land surface area range from ~0.2% to 0.4% (Burton-Johnson, Black, Fretwell, & Kaluza-Gilbert, 2016; Terauds et al., 2012), with glacial models suggesting that most if not all of this area has been covered by ice during multiple glacial cycles (DeConto & Pollard, 2016; Pollard & DeConto, 2016; Kaluza-Gilbert, 2016; Terauds et al., 2012), extending far back in time from hundreds of thousands to multi-million-year time-scales. While this evidence has been obtained far back in time from hundreds of thousands to multi-million-year time-scales, little is known about the timing of dispersal to this environment. This has led to a widely held view that most of the contemporary Antarctic biota must be of recent (post-LGM) origin (Convey et al., 2008).

Recent biological research has challenged this view, revealing many examples of species with long-term pre-glacial persistence. Examples can be found within many major groups of Antarctic terrestrial biota (e.g. invertebrates, lichens, mosses, diatoms and microbial groups; Allegrucci, Carchini, Todisco, Convey, & Sbordoni, 2006; Bennett, Hogg, Adams, & Hebert, 2016; Biersma et al., 2017; Biersma, Jackson, Stech, et al., 2018; Chong, Pearce, & Convey, 2015; Convey et al., 2008, 2009; Convey & Stevens, 2007; De Wever et al., 2009; Iakovenko et al., 2015; Pisa et al., 2014; Vyverman et al., 2010), extending far back in time from hundreds of thousands to multi-million-year time-scales. While this evidence has led to a paradigm shift in the perception of the age of Antarctic life, at present, biological and glaciological evidence still does not align and continues to challenge our understanding of the glacial history of Antarctica (Convey et al., 2008; Convey & Stevens, 2007).

The flora of Antarctica has a low species richness, and includes just c. 112 species of mosses (Ochyra, Smith, & Bednarek-Ochyra, 2008), c. 27 species of liverworts (Bednarek-Ochyra, Vana, Ochyra, & Smith, 2000) and two species of native angiosperms, the Antarctic pearlwort Colobanthus quitensis (Kunth.) Bartl. (Caryophyllaceae) and the Antarctic hair grass Deschampsia antarctica Desv. (Poaceae). Recent molecular research has revealed the Antarctic bryophyte flora to comprise a mixture of long-term survivors (Biersma et al., 2017; Biersma, Jackson, Stech, et al., 2018; Ochyra, 2003; Pisa et al., 2014) and more recent arrivals (Biersma, Jackson, Bracegirdle, et al., 2018; Biersma et al., 2017; Kato, Arikawa, Imura, & Kanda, 2013). While long-term survivors can be found within the bryoflora, the low diversity within the vascular flora suggests that it may be of recent origin. Fasanella, Premoli, Urdampilleta, González, and Chiapella (2017), studying the genetic diversity within D. antarctica, recently detected 17 nuclear DNA and six plastid DNA haplotypes in Patagonia, while Antarctica had just one nuclear and four plastid DNA haplotypes. As the haplotypes present in Antarctica were only a small fraction of those present in Patagonia, and the nuclear haplotype in Antarctica was also found in Patagonia, this suggested that the species likely dispersed to the Antarctic in the mid- to late-Pleistocene.

Although the genetic diversity of C. quitensis has previously been studied (Acuña-Rodríguez, Oses, Cortés-Vasquez, Torres-Díaz, & Molina-Montenegro, 2014; Androsiuk, Chwedorzewska, Szandar, & Giefwanowska, 2015; Cuba-Díaz, Cerda, Rivera, & Gómez, 2017; Cuba-Díaz, Klages, et al., 2017; Gianoli et al., 2004; Koc et al., 2018; Lee & Postle, 1975; Parnikoza, Maidanuk, & Kozeretska, 2007), as yet, no clear conclusions can be drawn about the age of the species in Antarctica (Parnikoza, Kozeretska, & Kunakh, 2011). This is mainly due to logistical and technical constraints, such as restricted geographical sampling and small sample sizes, and the genetic markers used being unsuitable for molecular dating techniques. Studies with more thorough sampling across the species’ biogeographic range and the use of more appropriate, DNA sequence-based markers are hence required to assess the timing of divergence among populations on either side of the Southern Ocean.

Here, by applying population genetic and molecular dating analyses to C. quitensis specimens collected from across the widest range of localities sampled to date, we aimed to assess (a) whether C. quitensis may have survived the LGM in refugia in the Maritime Antarctic (encompassing the Antarctic Peninsula, South Shetland Islands and South Orkney Islands), or (b) whether its arrival in these regions is a more recent post-glacial event.
2 MATERIALS AND METHODS

2.1 Sampling

The full biogeographic range of *C. quitensis* includes areas of the Antarctic Peninsula north from 69° S (Convey, Hopkins, Roberts, & Tyler, 2011), the South Shetland Islands, the South Orkney Islands, South Georgia, the Falkland Islands, the southern ranges of Chile and Argentina, the High Andes regions of Chile, Argentina, Ecuador and Bolivia, and extends into Mexico (Moore, 1970). Our dataset consisted of 270 samples collected from across the southern part of the species’ biogeographic range, where it is most commonly found (see Figure 1a; for the full distribution of the species see Figure 1b). To allow for a detailed study of both within-population and wider geographical variation, we combined two types of available datasets: (a) a population level dataset of a total of 200 freshly collected samples from 19 different field locations, with several (n > 1) samples collected per location, and (b) a dataset derived from single samples (n = 1) from 70 locations, derived from herbarium specimens, single fresh collections and one previously sequenced specimen from GenBank (see Table S1, Appendix S1). We included as many samples from the two datasets as possible in all analyses.

For the phylogenetic analyses, we included 21 samples (four from GenBank and 17 newly sequenced samples) from seven additional *Colobanthus* species as outgroups, viz., *C. subalatus* (D’Urv.) Hook.f., *C. kerguelensis* Hook.f., *C. apetalus* (Labill.) Druce, *C. strictus* Cheesem., *C. hookeri* Cheesem., *C. affinis* (Hook.) Hook.f. and *C. masonae* L.B.Moore. For the molecular dating analyses, nine specimens from *Sagina* species (seven from GenBank and two newly sequenced samples) were included as outgroups, based on the relationships reported by Dillenberger and Kadereit (2014) and Greenberg and Donoghue (2011) (see Table S1 for information on samples and GenBank accession numbers). Herbarium samples were obtained from the herbaria of the British Antarctic Survey, UK, and the University of Magallanes in Punta Arenas, Chile (herbarium codes AAS and HIP, respectively).

2.2 DNA extraction, PCR amplification, sequencing and alignment

The DNA regions selected for comparison included one nuclear ribosomal (nrDNA) marker, the ribosomal Internal Transcribed Spacer (ITS) region (ITS1-5.8S-ITS2) and four chloroplast (cpDNA) markers, viz., the *ndhF-rpl32* spacer, *rpl32-trnL* spacer, *trnQ-rps16* spacer and the *atpB-rbcL* spacer. For the first dataset (a), comprising only field-fresh collections, *ndhF-rpl32*, *rpl32-trnL* and *ITS* were sequenced, while for the second dataset (b), all markers were sequenced (see Table S1 for sampling locations and sequence details).

DNA was extracted from leaf tissue using the DNeasy Plant Mini Kit (Qiagen GmbH, Hilden, Germany), and E.Z.N.A. Plant DNA Kit (Omega Biotek, USA) following the manufacturers’ instructions, using liquid nitrogen and a mortar and pestle for tissue disruption. PCR amplification was carried out using the Taq PCR Core Kit (Qiagen GmbH, Hilden, Germany) and Platinum Taq DNA polymerase (Invitrogen, Life Technologies) according to the manufacturers’ instructions, with the addition of 1 μl of bovine serum albumin. Primer information and annealing temperatures are given in Table S2 (Appendix S1). Forward and reverse sequencing was performed by LGC Genomics (Berlin, Germany) and Macrogen (Seoul, Korea).

Forward and reverse sequences were combined and aligned with *prank* 140603 (Löytynoja & Goldman, 2008), using default settings, with minor corrections made manually. Models of DNA sequence evolution were selected using *jModelTest* 2.1.10 (Darriba, Taboada, Doallo, & Posada, 2012; Guindon & Gascuel, 2003) implementing the SPR base tree search, G rate variation option and the corrected Akaike information criterion (AICc) method for model comparisons. This found the most appropriate models were JC for *ITS*, TPM1uf for *atpB-rbcL*, and TPM1uf+G for *ndhF-rpl32*, *rpl32-trnL* and *trnQ-rps16*.

![FIGURE 1](image-url) (a) Map with sampling locations of *Colobanthus quitensis*, comprising locations of freshly collected samples from field sites (stars; with multiple samples per site) and samples from herbaria (circles; representing a single sample per location). Biogeographical regions are indicated with different colours. (b) Global distribution of *C. quitensis* (black dots representing the distribution of *C. quitensis*, following Moore, 1970; red dot representing southern limit of *C. quitensis*, following Convey et al., 2011).
Bayesian analyses were performed using MrBayes 3.2 (Ronquist et al., 2012). All analyses were run for $25 \times 10^6$ generations, applying default settings and the closest match to the JModelTest identified substitution models per partition (cpDNA: nst = 6, rates = gamma; ITS: nst = 1, rates = equal), sampling every $1.0 \times 10^3$ generations, and omitting the first 25% of trees as burn-in. Convergence was assessed using Tracer 1.6 (Rambaut, Suchard, Xie, & Drummond, 2014) by verifying that split frequencies had an average standard deviation of < 0.01 and all posterior parameter estimates exceeded effective sample sizes by >200. Maximum clade credibility trees with median heights were visualized using FigTree 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/). Maximum likelihood analyses were performed using RAxML-GUI 1.3.1 (Silvestro & Michalak, 2012), applying the ‘bootstrap+consensus’ option (1,000 iterations) using the JModelTest identified models of evolution with other settings as default. We inferred trees for ITS, trnQ-rps16, atpB-rbcL and the combined chloroplast regions ndhF-rpl32R and rpl32F-trnL (the latter were combined because they were always present), as well as generating a combined tree using ndhF-rpl32R, rpl32F-trnL and ITS (for which data from most specimens were included; n = 232). To assess for topological incongruence among phylogenies derived from the cpDNA and nrDNA partitions, we used >70% bootstrap (BS) and >95% posterior probability (PP) thresholds. Topological conflicts were assumed to be significant if two conflicting relationships for the same set of taxa were both supported with bootstrap values >70% and PP >95%. Phylogenetic analyses of the combined cpDNA and nrDNA datasets were conducted using an alignment containing unique sequences only (for a list of unique sequences see Table S1), that were extracted from the full dataset using Geneious 9.1.8 (https://www.geneious.com).

To assess within-species variation in C. quitensis according to biogeographic region, tcs phylogenetic networks (Templetion, Crandall, & Sing, 1992) were built using PopART (Leigh & Bryant, 2015), with default settings. Networks were made for each marker separately (ITS, trnQ-rps16, atpB-rbcL and the combined chloroplast regions ndhF-rpl32R and rpl32F-trnL) and from a combined dataset containing ndhF-rpl32R, rpl32F-trnL and ITS sequences. We calculated standard genetic diversity indices for all markers in Arlequin 3.5.1.2 (Excoffier & Lischer, 2010). We additionally carried out Tajima’s D (Tajima, 1989) and Fu’s F$_i$ (Fu, 1997) neutrality tests for the cpDNA regions only. For the combined ndhF-rpl32R and rpl32F-trnL dataset, we also calculated molecular diversity indices for biogeographic regions with sample sizes of >10 (High Andes, Patagonia, South Georgia, South Shetland Islands and Antarctic Peninsula). Additionally, pairwise F$_{ST}$ and $\Phi_{ST}$ (Excoffier, Smouse, & Quattro, 1992) values (using Kimura 2P genetic distances; Kimura, 1980) were calculated between these biogeographic regions, with 10,000 dataset permutations to assess significance. Numbers of variable and parsimony informative (PI) sites were calculated using MEGA7 (Kumar, Stecher, & Tamura, 2016).

### 2.4 Molecular dating

Relative divergence times and ages for C. quitensis were calculated using StarBEAST 2.5.1 (Bouckaert et al., 2014) on the combined ndhF-rpl32R, rpl32F-trnL and ITS dataset that also included nine Sagina specimens as outgroups. Analyses were performed using the unique haplotypes only. As there are no fossil or geological calibration points available for molecular dating within the genus, we used two alternative methods for date calibration: (a) employing a previously calculated divergence date for the split between Colobanthus and Sagina from Dillenberger and Kadereit (2017), in the form of a lognormal prior of 3.44 Ma and a 95% highest posterior density interval of 1.34–5.91 Ma, and (b) applying a substitution rate on the cpDNA partition of 0.8 ± 0.06 × 10$^{-9}$ subst./site/year, based on the rate estimated for chloroplast noncoding regions by Yamane, Yano, and Kawahara (2006) and previously applied on the Caryophyllaceae species Silene acaulis (Gussarova et al., 2015). For both methods, we applied a coalescent Bayesian Skyline tree prior, strict molecular clock and the JC69 and GTR+G models of evolution for ITS and cpDNA markers, respectively, allowing independent clocks for both genomic partitions. We used a linear multi-species coalescent with constant root as well as the appropriate ploidy level gene trees for both partitions. All runs had a chain length of $1.0 \times 10^8$ generations, logging parameters every $5.0 \times 10^3$ generations. Convergence was assessed in Tracer as described above. A maximum clade credibility tree with median node heights and 10% burn-in was constructed using TreeAnnotator 2.5.1 (Bouckaert et al., 2014) and visualized using FigTree 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/).
while these specimens formed distinct haplotypes in the shared with South American specimens within the specimens from the southern Maritime Antarctic fell within a group haplotype shared with South American specimens. Conversely, work, most of these specimens fell within a common sequence from the South Shetland Islands plus northern Antarctic Peninsula southern Antarctic Peninsula and the other containing specimens fell into two distinct groupings: one containing specimens from the Maritime Antarctic (Figure 3c) revealed that specimens from the Maritime Antarctic 

| Marker               | n  | bp* | v  | PI | π    | h    | Tajima’s D (p) | Fu’s F_s (p) |
|---------------------|----|-----|----|----|------|------|---------------|--------------|
| (a)                 |    |     |    |    |      |      |               |              |
| ITS                 | 263| 536 | 8  | 7  | 0.005 ± 0.003 | 0.771 ± 0.025 | N.A.          | N.A.         |
| trnQ-rps16         | 79 | 712 | 6  | 3  | 0.003 ± 0.002 | 0.604 ± 0.076 | −1.059 (.147) | −1.495 (.279) |
| atpB-rbcL          | 66 | 822 | 8  | 5  | 0.002 ± 0.001 | 0.941 ± 0.033 | −1.864 (.006)* | −6.627 (.001)* |
| ndhF-rpl32+trnL    | 226| 979 | 28 | 16 | 0.009 ± 0.004 | 0.837 ± 0.023 | −1.552 (.030)* | −3.494 (.251) |
| (b)                 |    |     |    |    |      |      |               |              |
| High Andes         | 52 | 923 | 6  | 5  | 0.007 ± 0.004 | 0.799 ± 0.023 | N.A.          | N.A.         |
| Patagonia           | 75 | 968 | 22 | 11 | 0.004 ± 0.002 | 0.915 ± 0.016 | N.A.          | N.A.         |
| South Georgia       | 16 | 913 | 2  | 1  | 0.001 ± 0.001 | 0.617 ± 0.135 | N.A.          | N.A.         |
| S. Shetland Is.    | 89 | 913 | 0  | 0  | 0.000 ± 0.000 | 0.044 ± 0.030 | N.A.          | N.A.         |
| Ant. Peninsula      | 29 | 913 | 2  | 2  | 0.001 ± 0.001 | 0.687 ± 0.050 | N.A.          | N.A.         |

Abbreviations: bp*: no. of usable base pairs (loci < 5.0% missing data); For Tajima’s D and Fu’s F_s neutrality tests p < .05 is significant (*); h: gene diversity; n: number of samples; PI: parsimony informative sites; v: variable sites; π: nucleotide diversity (average over locus).

regions (Figure 2; see Figure S1.1 and S1.2a–d in Appendix S1 for phylogenetic trees of combined and single markers, respectively). Phylogenetic analyses of C. quitensis indicated a north-to-south expansion, with an early split of several High Andes populations from the remaining biogeographic regions (Figure 2). Antarctic haplotypes were associated with two different clades: one clade contained specimens from the northern Maritime Antarctic, South Shetland Islands and South Georgia, while the other clade consisted of a polytomy containing specimens from the southern Antarctic Peninsula as well as many other biogeographic regions. Other clades contained High Andes samples from the southernmost Andes location (La Parva; see Table S1), and Patagonian specimens, respectively.

For the population genetic analyses of ITS, the 263 individuals analysed yielded nine unique haplotypes, while for the combined rpl32-trnL+ndhF-rpl32 regions, the 226 individuals analysed yielded 28 haplotypes. The 79 and 66 individuals analysed for trnQ-rps16 and atpB-rbcL both resulted in seven haplotypes. The TCS networks of ITS (Figure 3a) and rpl32-trnL+ndhF-rpl32 (Figure 3b) both revealed that the distribution of Antarctic specimens fell among different groups: in the ITS network, one of these consisted solely of specimens from the northern Maritime Antarctic, South Shetland Islands and South Georgia, while in the rpl32-trnL+ndhF-rpl32 network, most of these specimens fell within a common sequence haplotype shared with South American specimens. Conversely, specimens from the southern Maritime Antarctic fell within a group shared with South American specimens within the ITS network, while these specimens formed distinct haplotypes in the rpl32-trnL+ndhF-rpl32 network. The combined analysis of these markers (Figure 3c) revealed that specimens from the Maritime Antarctic fell into two distinct groupings: one containing specimens from the southern Antarctic Peninsula and the other containing specimens from the South Shetland Islands plus northern Antarctic Peninsula (Figure 3d). Both groupings also contained one or several specimens from South Georgia, respectively. The trnQ-rps16 network (Figure 3e) also showed two distinct haplotypes containing Antarctic specimens, however both were shared with South American specimens. The atpB-rbcL network (Figure 3f) showed only one haplotype containing Antarctic specimens, which was shared with South American specimens.

All pairwise $F_{ST}$ and Φ_{ST} comparisons between biogeographic regions were significant (Table 2), with the South Shetland Islands showing particularly high population differentiation in haplotypic diversity ($F_{ST}$), followed by South Georgia and the Antarctic Peninsula. Taking into account molecular distances ($Φ_{ST}$), the High Andes populations were particularly differentiated from most other regions.

### 3.3 Divergence time analysis

The estimated age using the fossil-calibrated method (I) revealed an earlier split of the most recent common ancestor (TMRCA: split Sagina - Colobanthus) than the rate-informed dating analysis (II) (Table 3). Both methods suggested that the genus Colobanthus diverged throughout the course of the Pleistocene, and that C. quitensis originated c. 0.181 (0.042–0.431) to 0.666 (0.401–0.999) Ma (method I and II, respectively; i.e. the calculated age of the root of C. quitensis, corresponding to the split between the majority of the High Andes populations and the remaining populations). Further date estimates for divergences of other populations within this young species were not possible due to a lack of sufficient variation within the sequenced DNA regions. Rates for both partitions calculated using method (I) were $7.20 ± 0.06 × 10^{-9} \text{subst./site/year for ITS, and}$ $2.27 ± 0.06 × 10^{-8} \text{subst./site/year for ndhF-rpl32+trnL}$. The rate calculated for ITS using method (II) was approximately sixfold lower, at $1.13 ± 0.01 × 10^{-9} \text{subst./site/year}$. 

| Marker               | n  | bp* | v  | PI | π    | h    | Tajima’s D (p) | Fu’s F_s (p) |
|---------------------|----|-----|----|----|------|------|---------------|--------------|
| ITS                 | 263| 536 | 8  | 7  | 0.005 ± 0.003 | 0.771 ± 0.025 | N.A.          | N.A.         |
| trnQ-rps16         | 79 | 712 | 6  | 3  | 0.003 ± 0.002 | 0.604 ± 0.076 | −1.059 (.147) | −1.495 (.279) |
| atpB-rbcL          | 66 | 822 | 8  | 5  | 0.002 ± 0.001 | 0.941 ± 0.033 | −1.864 (.006)* | −6.627 (.001)* |
| ndhF-rpl32+trnL    | 226| 979 | 28 | 16 | 0.009 ± 0.004 | 0.837 ± 0.023 | −1.552 (.030)* | −3.494 (.251) |

Abbreviations: bp*: no. of usable base pairs (loci < 5.0% missing data); For Tajima’s D and Fu’s F_s neutrality tests p < .05 is significant (*); h: gene diversity; n: number of samples; PI: parsimony informative sites; v: variable sites; π: nucleotide diversity (average over locus).
4 | DISCUSSION

4.1 | The origin of *Colobanthus quitensis* in Maritime Antarctica and in South Georgia

Most genetic markers revealed multiple alleles within the Maritime Antarctic (Antarctic Peninsula, South Shetland Islands and South Orkney Islands) that were shared with regions further north (all markers except atpB-rbcL; Figure 3), suggesting *C. quitensis* dispersed to the region at least twice, once to the northern Antarctic Peninsula and South Shetland Islands, and once to the southern Antarctic Peninsula (Figure 3d). Both Maritime Antarctic regions shared identical haplotypes with populations from South Georgia, suggesting that these regions, physically separated by ~850–1,300 km, are more closely connected than has been previously thought. The direction in which dispersal events have taken place is not clear. As there is a South Georgian sample in the centre of the TCS network (Figure 3c), it is possible that one or both Maritime Antarctic groups dispersed from South Georgia. However, it is also plausible that the species dispersed from the Maritime Antarctic to South Georgia, especially given the general direction of oceanic and atmospheric currents that characterize this region (Biersma, Jackson, Bracegirdle, et al., 2018). A shared haplotype between populations in southern South America (Patagonia and Falkland Islands) and South Georgia (Figure 3c) suggests that a recent dispersal event from southern South America to South Georgia has also occurred.

Both Maritime Antarctic and South Georgia groups were most closely related (being only one mutational step separated in the compiled cpDNA and nrDNA TCS network; Figure 3c) to the main haplotype comprising Patagonia, the Falkland Islands and South Georgia, and could therefore have originated from any of these latter regions. As both Maritime Antarctic populations share identical haplotypes with South Georgia, and as there are suggestions that the latter may have harboured LGM ice-free refugia (Allegrucci et al., 2006; McCracken, Wilson, Peters, Winker, & Martin, 2013; Van der Putten, Verbruggen, Ochyra, Verleyen, & Frenot, 2010), this location could have been a potential source and refugium for one or both of the Maritime Antarctic populations. This possibility is, however, counter to the general direction of oceanic and atmospheric currents noted above. Alternatively, southern South America could also have been the original source location, with this region being thought to have harboured various Pleistocene
refugia (Sersic et al., 2011). Intriguingly, the southern Maritime Antarctic group also showed a close affinity to the northern High Andes populations (separated by only two mutational steps; Figure 3b,c), a finding worthy of investigation in future studies.

4.2 Recent arrival of the Antarctic vascular flora

Exactly when the dispersal events across the Southern Ocean occurred is not certain, but the shared haplotypes and genotypes with specimens from South Georgia as well as the genetic similarity to specimens from South American regions suggest that C. quitensis reached the Antarctic on a relatively recent (late-Pleistocene) timescale. As we report here, previous studies have recorded low genetic variation within C. quitensis (Acuña-Rodríguez et al., 2014; Androsiuk et al., 2015; Koc et al., 2018; Lee & Postle, 1975), suggesting a recent spread of the species across the Antarctic Peninsula. This stands in contrast with an earlier suggestion that C. quitensis is a likely pre-glacial relict present in Antarctica since the Oligocene–Pliocene (Parnikoza et al., 2007). Notably, we also find that C. quitensis is itself a relatively young species (<1 Ma; see Table 3), and much younger than the Oligocene-Pliocene. Our overall results suggest that C. quitensis likely only became established in the Maritime Antarctic on a late-Pleistocene timescale, and, although we cannot be certain about its exact arrival time, it possibly only arrived there after the initial post-LGM ice retreat in the Antarctic Peninsula and South Shetland Island regions (c. 12–14 ka; Anderson, 2002, and references therein).
inference of multiple successful colonizations of a vascular plant species to the Antarctic over a relatively short time-scale (since late-Pleistocene) is of significance with respect to predicting future colonizations of vascular plants and other organisms on the Antarctic Peninsula, in particular with the increase of ice-free land associated with regional warming (Lee et al., 2017) and recent human activity in the area (Convey & Peck, 2019).

This study fills a significant gap in knowledge of the origin of the Antarctic terrestrial flora. The other native angiosperm, the grass *D. antarctica*, has been the subject of more population genetic studies than *C. quitensis*. For example, Van de Wouw, Dijk, and Huiskes (2008), using amplified fragment length polymorphisms (AFLPs) and chloroplast sequences, detected a low genetic diversity in this grass in the Antarctic, suggesting it was unlikely that *D. antarctica* survived the LGM in Antarctica *in situ*. Subsequently, Fasanella et al. (2017), studying patterns of genetic variability of *D. antarctica* within populations from across both sides of the Drake Passage, identified eight chloroplast haplotypes, of which Antarctic populations included four haplotypes (one unique, the remaining overlapping with Patagonian haplotypes). In the more variable nuclear marker (ITS), 17 haplotypes were found in total, of which Antarctic populations included only one haplotype, which was also present in Patagonia. Overall, the results suggested a mid- to late-Pleistocene arrival of the grass in Antarctica, corroborating our findings for *C. quitensis*. Future studies with new markers (such as those identified by Ishchenko, Panchuk, Andreev, Kunakh, & Volkov, 2018; Rabokon et al., 2017; Biersma, Jackson, Stech, et al., 2018; Ochyra, 2003; Pisa et al., 2014) and lichens (including many endemic species; Green, Sancho, Türk, Seppelt, & Hogg, 2011; Øvstedal & Smith, 2001), the likely late-Pleistocene arrival of the Antarctic vascular flora is therefore a notable exception to this generalization.

### Table 3

| Method | TMRCA Sagina – Colobanthus | TMRCA Colobanthus | TMRCA C. quitensis |
|--------|----------------------------|-------------------|-------------------|
| I      | 3.236 (1.341–5.850) Ma     | 0.408 (0.112–0.896) Ma | 0.181 (0.042–0.431) Ma |
| II     | 1.876 (0.911–3.307) Ma     | 1.450 (0.847–2.137) Ma | 0.666 (0.401–0.999) Ma |

*Based on a previously calculated divergence date for the split between Colobanthus and Sagina (Dillenberger & Kadereit, 2017).

*Based on estimated substitution rate for noncoding chloroplast regions (Yamane et al., 2006).

### 4.3 Genetic variation of *Colobanthus quitensis* within southern South America

Based on the sampling included in this study, an early split could be found between the majority of the populations from the central South American Andes and those from the remaining populations, including Patagonia (Figure 2). We note that we did not have access to material from populations from areas further north in South America and in southern North America (Mexico), where the species is also sporadically found (see Figure 1b). The high genetic variation and abundance of this species in Patagonia (Table 1; Figure 3a–c) suggests that populations have persisted and remained stable in this region for a long period, and may indicate possible presence within multiple refugia during the Pleistocene, as also found for other species (Sersic et al., 2011).

The southernmost High Andes population sampled (La Parva, near Santiago, Chile) had nine specimens that converged with other more northern High Andes populations, but also five specimens that grouped within the polytomy containing all southern populations (Figure 2). In the haplotype network (Figure 3a), these five samples were equally closely related to southern South American haplotypes and the southern Maritime Antarctic group. The observation that this southern High Andes population at La Parva shares haplotypes with other High Andes populations, as well as with more southerly populations, suggests that there is genetic admixture between the High Andes and Patagonian populations in this region.

With its origin in the Andean range and/or cold regions of Patagonia (see Figure 2), *C. quitensis* is likely pre-adapted to cold, high altitude environments, which are characterized by highly variable conditions (e.g. in temperatures and water availability). The genetic similarity of *C. quitensis* across its current biogeographical distribution suggests ecological niche conservatism in its ability to withstand harsh and/or variable conditions (as shown by its tolerance to cold, moderately saline and/or dry environments), combined with opportunistic dispersal capabilities to reach and colonize other suitable habitats (e.g. Antarctic and sub-Antarctic environments). Overall, the genetic information shown here may be useful for future studies that apply niche comparative methods to link macroclimatic variables in explaining the past, present and future distribution of *C. quitensis*.

### 4.4 Dispersal within the genus Colobanthus

Distribution patterns within *Colobanthus* suggest that the genus is efficient at dispersing to other regions, including across oceans. The outgroup species *C. subulatus* showed identical sequences in samples from Patagonia and South Georgia in all markers (Figure S1.2a–d), suggesting that this species has also recently arrived in South Georgia from Patagonia. Similarly, in *C. kerguelensis*, nearly identical sequences in all markers were found across the remote Kerguelen Islands, Crozet Islands and Amsterdam Island in the Indian Ocean.
While the overall distribution of Colobanthus appears Gondwanan (including representatives from New Zealand, Australia, South America, many sub-Antarctic islands and Antarctica), the genus’ age is clearly much younger than the break-up of Gondwana (see Table 3 and Dillenberger & Kadereit, 2017), supporting the hypothesis that many Colobanthus species are efficient trans-oceanic dispersers. This is confirmed by their presence on various geologically young islands, such as sub-Antarctic Prince Edward Island (c. 215 ka; Hänel & Chown, 1998). Further phylogeographical studies are required to assess historical dispersal and speciation patterns within the wider genus.

4.5 Possible modes of dispersal

While bryophytes and other spore-dispersed biota could have been distributed to Antarctica by wind (e.g. see Biersma, Jackson, Bracegirdle, et al., 2018), the weight of the seeds of C. quitensis (~50 μg) probably prevents such dispersal. Oceanic dispersal (‘rafting’) or animal vectors (e.g. migrating birds) are both more likely routes by which the species could have arrived in Antarctica. The species is known to be moderately salt-tolerant, and in Patagonia, South Georgia and the Maritime Antarctic commonly occurs in many coastal environments, including the top of the intertidal zone in Patagonia (Cuba-Díaz, Castel, Acuña, Machuca, & Cid, 2017). Whether its seeds could survive exposure to seawater during rafting is unknown, but recently an example of rafting kelp has revealed that Antarctica is not completely isolated from biological sea-rafting particles from mid-latitude source populations (Avila et al., 2020).

Another possible mode of dispersal for C. quitensis could have been via the plumage of common Antarctic birds, such as gulls (Parnikoza et al., 2012, 2018). However, C. quitensis seeds are smooth and have no hooks or spines to facilitate their attachment to bird plumage, lessening the likelihood of this type of trans-oceanic dispersal. Alternatively, dispersal via the guts of birds, such as the white-rumped sandpiper (Calidris fuscicollis) could have facilitated a historical dispersal event. This long-distance migrating shorebird, breeding in the North American Arctic and wintering in southern South America and the Falkland Islands, is also a rare visitor to South Georgia and the South Shetland Islands (Trivelpiece et al., 1987), where sightings have increased over the last 30 years (Korczak-Abshire, Angiel, & Wierzbicki, 2011). The species has also been observed east of the Andes for biogeographical history. (Polish Polar Research, 29, 320–326).

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DATA AVAILABILITY STATEMENT

Sequence data have been submitted to the GenBank database under accession numbers MN640112–MN640391 and MN614479–MN615128 (see Table S1, Appendix S1). Phylogenetic and Popart matrixes are available in the Dryad Digital Repository (https://doi.org/10.5061/dryad.qrjf6q5bw).

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**BIOSKETCH**

Elisabeth Machteld Biersma is an evolutionary biologist at the British Antarctic Survey, studying the distribution and origin of Antarctic biota using population genetics and molecular dating techniques. This project was a joint research effort with Cristian Torres-Díaz to develop a large-scale dataset of *C. quitensis*.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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