Inter-island dispersal of flightless *Bothrometopus huntleyi* (Coleoptera: Curculionidae) from the sub-Antarctic Prince Edward Island archipelago

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**Abstract:** *Bothrometopus huntleyi* is a flightless weevil endemic to the volcanically-formed sub-Antarctic Prince Edward Islands archipelago that arose approximately 0.5 million years ago (m.y.a.). Since emergence, a series of volcanic and glaciation events have occurred on Marion Island, whilst Prince Edward Island, the second island constituting the archipelago, has remained largely unaffected by glaciation. Cytochrome oxidase I gene analyses indicate that major historical dispersal events in this species are linked to the geologically discrete histories of these islands and underlie the high haplotype diversity (0.995) recovered for the Prince Edward Islands archipelago. The estimated time to haplotype coalescence of ~0.723 m.y.a. is in keeping with estimated dates of island emergence, and the majority of individuals appear to have descended from a relict, high-altitude population that is still present on Marion Island. The first major inter-island dispersal event occurred ~0.507 m.y.a., coinciding with the oldest dated rocks on Marion Island. Apart from this early inter-island colonization, only one other between-island dispersal event was detected. The genetically discrete *B. huntleyi* complexes on each of the islands of the Prince Edward Islands archipelago together with the low levels of inter-island gene flow reaffirm the need to control alien invasive mice, which are restricted to Marion Island, and which prey on this weevil species.

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**Key words:** COI gene, conservation biogeography, invasion biology, mtDNA, phylogeography

**Introduction**

*Bothrometopus huntleyi* (Grobler et al. 2011) (Coleoptera: Curculionidae) belongs to the *Ectemnorhinus* group of genera (Kuschel & Chown 1995), also known as the tribe Ectemnorhinini (Alonso-Zarazaga & Lyal 1999), which is restricted to islands of the sub-Antarctic South Indian Ocean Province (SIP). This group of weevils comprises eight genera and 37 flightless species (Kuschel & Chown 1995, Grobler et al. 2006, 2011), of which seven occur on the Prince Edward Islands. Of these seven species, five, including *B. huntleyi*, are endemic to the archipelago, with *Bothrometopus randi* (Jeannel 1953) also being found on the Iles Crozet to the east, and *Palirhoeus eatoni* (C.O. Waterhouse 1876) occurring on all SIP islands (Kuschel & Chown 1995). Recent molecular studies both of this group of weevils (Grobler et al. 2006) and of other taxa in the region (e.g. Stevens et al. 2006, Myburgh et al. 2007) have suggested that relationships among species, within and among islands and archipelagos (including the Prince Edward Island archipelago) may be much more complex than suggested by morphological analyses. In consequence, it may be argued that the status of *B. huntleyi* on the Prince Edward Island archipelago has not been fully resolved, and that it might be expected to show substantial phylogeographic structure associated with the different histories of the two islands which lie only 19 km apart. Perhaps most significantly, Marion Island, the larger of the two Prince Edward Islands, was extensively glaciated whilst the smaller Prince Edward Island shows no evidence of glaciation (Boelhouwers et al. 2008). McDougall et al. (2001) suggested that Marion Island had five distinct cold periods and eight volcanic ages, with two of these accompanying major glaciations (Fig. 1). Although the exact sequence and spatial extent of volcanic and glacial events is currently the subject of revision, this overall scenario appears to be robust (Boelhouwers et al. 2008).

In consequence, substantial differences among populations of *B. huntleyi* on the two islands might be expected. Such differences would not only be of biogeographic significance (see e.g. Chown 1992), but would also have substantial conservation implications given the differences in invasive alien species on the islands, especially the absence of house mice on Prince Edward Island, which have a dietary preference for weevils on Marion Island (Chown & Smith 1993). Here we set out to examine the evolutionary dynamics shaping the population structure of *B. huntleyi*, which occurs on both islands, is widely distributed across Marion Island, and has until recently been confounded with *B. parvulus* (Grobler et al. 2011).
Materials and methods

Systematics and biology

Although the genus Bothrometopus has previously been separated into two distinct groups, the fasciatus- and gracilipes-groups, which are distinguished by, respectively, the absence and presence of vaginal spicules (see Dreux & Voisin 1984, 1986, Kuschel & Chown 1995 for earlier work on the genus and its now synonymous allies), Grobler et al. (2011) have called for the re-evaluation of these groups.

Because B. huntleyi was previously not distinguished from the very similar B. parvulus, and because it is now clear that B. parvulus is restricted to coastal areas of both islands (Grobler et al. 2011), it can be reasonably assumed, from the previous literature on habitat use in the group (e.g. Chown 1989), that B. huntleyi predominantly occurs in epilithic moss cushions from coastal rock faces to high-altitude fellfield and polar desert. The larvae and adults feed on algae, lichens, and bryophytes (Chown 1989, 1992). Individuals may occur occasionally on Azorella selago, but feed on epiphytic algae and bryophytes, rather than on the plant itself (Chown 1989, 1992). On both Marion Island and Prince Edward Island, B. huntleyi shows distinct variation in size, associated both with elevation and habitat type, although differences among the two islands are not pronounced (Chown 1992, Chown & Smith 1993, Chown & Klok 2003).

Study sites and sampling

Specimens of B. huntleyi were collected during three consecutive years (April 2001–April 2003) from 14 localities from the eastern side of Marion Island (Fig. 2). Due to restricted access to Prince Edward Island, weevils were only sampled from five localities on one occasion in April 2003 (Fig. 2). For each island, the specimens were collected from a range of altitudes, and individuals selected for genetic characterization were chosen to ensure the broadest possible size and colour variation representation, for each sampling locality. The geographic coordinates, altitude and weevil size range recorded for each sampling locality are summarized in Table I. All specimens were collected by hand and preserved in absolute ethanol. Bothrometopus brevis (C.O. Waterhouse) from Heard Island was selected as an outgroup, since it is the closest known sister taxon of B. huntleyi (Grobler et al. 2011, Table I).

Genetic characterization

DNA from each individual was extracted from a single leg which, following removal from ethanol was washed and rehydrated in distilled water for 10 minutes, prior to being frozen in liquid nitrogen and ground using an Eppendorf micro pestle (Merck, South Africa). DNA was extracted using the High Pure PCR Template Preparation Kit (Roche Applied Science) using the supplier’s procedure for the isolation of nucleic acids from mammalian tissue with modification to the proteinase K tissue lysis incubation step, which was performed for 48 hr instead of the recommended 1 hr for mammalian tissue.

COI primers, GF-1858 and GR1-2938 which are specific for weevils from Marion Island (Grobler et al. 2006) were used to amplify a 1059 bp PCR product under previously described reaction conditions (Grobler et al. 2006) using a thermal cycling profile that consisted of an initial denaturation step at 94°C for 90 s, followed by 40 cycles of 94°C for 22 s,
Fig. 2. Map indicating flightless weevil (Bothrometopus huntleyi) sampling localities on the Prince Edward Island archipelago and that correspond to the geographic coordinates summarized in Table I.

Table I. Summary of the 19 sampling localities on Marion Island (MI) and Prince Edward Island (PEI) from which the genetically characterized Bothrometopus huntleyi specimens included in this study were collected.

<table>
<thead>
<tr>
<th>Island</th>
<th>Sampling locality (a.s.l.)</th>
<th>Geographic coordinates</th>
<th>Number of specimens per locality (size range in mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MI</td>
<td>Kildalkey Bay (0 m)</td>
<td>46°57'58.3&quot;S, 3°51'22.2&quot;E</td>
<td>1 (3.92)</td>
</tr>
<tr>
<td>MI</td>
<td>Ship's Cove (0 m)</td>
<td>46°51'41&quot;S, 37°50'66&quot;E</td>
<td>8 (3.31–5.00)</td>
</tr>
<tr>
<td>MI</td>
<td>Trypot Beach (0 m)</td>
<td>46°53'05.2&quot;S, 37°52'06&quot;E</td>
<td>1 (3.30)</td>
</tr>
<tr>
<td>MI</td>
<td>Stony Ridge (150 m)</td>
<td>46°54'88.1&quot;S, 37°51'48.4&quot;E</td>
<td>4 (4.61)</td>
</tr>
<tr>
<td>MI</td>
<td>Junior’s Kop (200 m)</td>
<td>46°52.794'S, 37°50.083'E</td>
<td>4 (3.15–5.00)</td>
</tr>
<tr>
<td>MI</td>
<td>Tafelberg (250 m)</td>
<td>46°53'03.5&quot;S, 37°48'20.1&quot;E</td>
<td>1 (3.69)</td>
</tr>
<tr>
<td>MI</td>
<td>Repetto’s Hill (300 m)</td>
<td>46°50'35.7&quot;S, 37°44'21.8&quot;E</td>
<td>4 (3.36–4.77)</td>
</tr>
<tr>
<td>MI</td>
<td>First Red Hill (400 m)</td>
<td>46°53.412'S, 37°48.21'E</td>
<td>5 (2.80–4.08)</td>
</tr>
<tr>
<td>MI</td>
<td>Long Ridge South (450 m)</td>
<td>46°52'45&quot;S, 37°47'00&quot;E</td>
<td>2 (3.08–5.08)</td>
</tr>
<tr>
<td>MI</td>
<td>First Red Hill (600 m)</td>
<td>46°53.647'S, 37°47.208'E</td>
<td>4 (2.96–4.72)</td>
</tr>
<tr>
<td>MI</td>
<td>Halfway Kop (600 m)</td>
<td>46°54'06&quot;S, 37°47'83&quot;E</td>
<td>6 (3.08–5.31)</td>
</tr>
<tr>
<td>MI</td>
<td>Feldmark Plateau (600 m)</td>
<td>46°56'35&quot;S, 37°46'10&quot;E</td>
<td>2 (3.39–5.15)</td>
</tr>
<tr>
<td>MI</td>
<td>Katedraalkrans (800 m)</td>
<td>46°53.896'S, 37°46.482'E</td>
<td>4 (3.60–4.85)</td>
</tr>
<tr>
<td>MI</td>
<td>Ice Plateau (1000 m)</td>
<td>46°54.29'S, 37°45.375'E</td>
<td>8 (3.92–5.69)</td>
</tr>
<tr>
<td>PEI</td>
<td>Cave Bay (0 m)</td>
<td>46°38.752'S, 37°59.780'E</td>
<td>4 (2.46–4.62)</td>
</tr>
<tr>
<td>PEI</td>
<td>(200 m)</td>
<td>46°38.457'S, 37°58.396'E</td>
<td>5 (3.31–4.39)</td>
</tr>
<tr>
<td>PEI</td>
<td>(400 m)</td>
<td>46°38.211'S, 37°57.482'E</td>
<td>5 (3.31–4.31)</td>
</tr>
<tr>
<td>PEI</td>
<td>(600 m)</td>
<td>46°37.533'S, 37°55.985'E</td>
<td>4 (3.54–5.00)</td>
</tr>
<tr>
<td>PEI</td>
<td>TvZB (672 m)</td>
<td>46°37.590'S, 37°55.891'E</td>
<td>4 (3.54–4.46)</td>
</tr>
</tbody>
</table>

a.s.l. = above sea level, TvZB = samples collected at the top of Van Zinderen Bakker.
46°C for 30 s and 72°C for 1 min and concluding with a final extension step of 1 min at 72°C. PCR products of the correct size were purified directly from the tube using a Roche High Pure PCR Product Purification Kit (Roche Applied Science). Automated cycle sequencing with the ABI PRISM Big Dye™ Terminator version 3.0 (Applied Biosystems) and two internal primers, GF5-1940 and GR5-2935 (Grobler et al. 2006), in separate reactions, was performed at an annealing temperature of 46°C. Sequences were viewed, edited and aligned in MEGA4 (Tamura et al. 2007), resulting in a homologous dataset 885 bp in length.

**Phylogenetic analyses**

Neighbor-Joining (NJ) and Minimum Evolution (ME) algorithms were used to construct distance trees in MEGA4 (Tamura et al. 2007) and maximum likelihood analyses were performed in PhyML (Guindon & Gascuel 2003) using the model of sequence evolution identified as the best-fit model under the Akaike Information Criterion (AIC) in ModelTest version 3.06 (Posada & Crandall 1998). Nodal support was assessed by 100 000 and 5000 bootstrap replications, respectively. Maximum parsimony (MP) analyses were performed in PAUP* v4.0b10 (Swofford 2003). Characters had equal weights in the initial analysis and were subsequently reweighted using the rescaled consistency index (RCI). Starting trees were obtained by random stepwise addition of sequences and branch swapping was performed using the tree-bisection-reconnection (TBR) algorithm. Bayesian Inference (BI) using MrBayes version 3.1 (Huelsenbeck & Ronquist 2001) was performed using priors guided by the best-fit model and parameters in ModelTest (Posada & Crandall 1998), namely the HKY85 model (Hasegawa et al. 1985) with the proportion of invariable sites (I) and gamma distribution shape parameter (G) being 0.791 and 1.6671, respectively. The analysis was initiated with random starting trees and run for 10 000 000 generations with Markov chains sampled every 2000 generations. Of the 5000 trees obtained, 1250 were discarded as “burn-in”. Tracer plots were visually inspected and tracer diagnostics (standard deviation of split frequencies, effective sample size), as implemented in MrBayes and Tracer v1.4 (Drummond & Rambaut 2007) were inspected to ensure that stationarity had been reached.

The equality of evolutionary rates between lineages was evaluated using the relative rate test (Li & Bousquet 1992) in PHYLYTEST version 2.0 (Kumar 1996). The likelihood ratio test (Felsenstein 1981, 1988) was also performed by calculating and comparing the log likelihood scores with and without the molecular clock enforced. Divergence times were estimated using the 2.3% nucleotide sequence divergence per million years, estimated from an arthropod mtDNA survey (Brower 1994) and the appropriate parameter-rich model of evolution (Papadopoulou et al. 2010) selected under the AIC in ModelTest (Posada & Crandall 1998). BEAST 1.5.3 (Drummond & Rambaut 2007) was used to obtain an ultrametric tree using Bayesian MCMC analysis orientated towards rooted, time-measured phylogenetics. Well supported nodes identified following NJ, ML and BI analyses were constrained to be monophyletic and the HKY+1+Γ model was enforced using a strict molecular clock model. The results of two independent runs were merged and analysed with Tracer v1.4 and TreeAnnotator v1.4.7 (Drummond & Rambaut 2007).

The program TCS version 1.21 (Clement et al. 2000) was used to generate a haplotype cladogram displaying the number of base pair differences between haplotypes. TCS version 1.21 (Clement et al. 2000) incorporates the cladogram estimation algorithm described by Templeton et al. (1992) and provides 95% parsimoniously plausible branch connections between the different haplotypes.

**Mismatch analyses, population expansion and diversity estimates**

Pairwise mismatch distributions, where the observed pairwise mismatch distributions were fitted to a stepwise expansion model by a generalized least square procedure following Schneider & Excoffier (1999), as implemented in Arlequin version 2.0 (Schneider et al. 2000), were generated. A constant size population is expected to show a ragged, multimodal distribution, while an expanding population is consistent with a smooth unimodal distribution. In each case, the raggedness index assesses the match of the real data to the model while the overall validity of the estimated expansion model was tested by comparing the distribution of the test statistic, SSD (sum of squared differences) between the observed and the estimated mismatch distribution using a bootstrap approach. Evidence for departure from the estimated expansion model is given by significant SSD values (Excoffier & Schneider 1999).

The validity of a stepwise expansion model for the data was tested by Monte Carlo Markov chain simulations (1000 steps) in Arlequin version 2.0 (Schneider et al. 2000). The time of the main expansion in generations (t) was estimated from the equation $\tau = 2\mu t$ using the moment estimator of time to the expansion ($\tau$) computed in Arlequin version 2.0 (Schneider et al. 2000) and a mutation rate ($\mu$) of 2.3% nucleotide sequence divergence per million years for arthropod mtDNA (Brower 1994). As the weevils complete one generation in one-year (Chown & Scholtz 1989), t could also be used as the time of the main expansion in years. Arlequin version 2.0 (Schneider et al. 2000) was utilized to estimate the essential population parameter $\theta$ using Watterson’s (1975) estimate, which is based on the number of segregating sites among the sequences, and Tajima’s (1983) estimate, which is based on the calculation of the mean number of pairwise differences of the sequences. Tajima’s estimate of $\theta$ puts more weight on ancient mutations and therefore reflects ancient population events (Fu 1997).
Fig. 3. Neighbour-Joining (NJ) tree of Bothrometopus huntleyi from Marion Island (MI) and Prince Edward Island (PEI) and B. brevis outgroup from Heard Island, based on 885 nucleotides of the mitochondrial cytochrome oxidase I (COI) gene inferred using the HKY85 model of sequence evolution. The taxon name contains the species designation, sample number, body length measurement, altitude and/or sampling locality, sex and island of origin. Nodal support values $\geq 60$ obtained from NJ and ML (100 000 and 5000 bootstrap replications, respectively) and $\geq 90$ from Bayesian Inference (BI) and expressed as a percentage, are indicated NJ/ML/BI. ‘- -’ denotes bootstrap support values below 60 (NJ and ML) and below 90 (BI).
Fu’s (1997) $F_S$ test of neutrality, as estimated in Arlequin version 2.0 (Schneider et al. 2000), and the $R_2$ test of neutrality (Ramos-Onsins & Rozas 2002), as estimated in DNASP version 5 (Librado & Rozas 2009), were shown to be powerful tests for detecting recent population expansions under assumptions of neutrality with $R_2$ showing better results for small sample sizes and $F_S$ for large sample sizes. The $R_2$ statistic is based on the difference between the number of singleton mutations and the average number of nucleotide differences among sequences within a population sample (Ramos-Onsins & Rozas 2002). The haplotype diversity ($h$) and nucleotide diversity ($\pi$) were estimated in DNASP version 5 (Librado & Rozas 2009).

Results

A homologous region of 885 bp corresponding to the 5′ end of the COI gene was generated for 73 $B. huntleyi$ individuals and two $B. brevis$ outgroup specimens. All sequences have been deposited in the Genbank database under accession numbers GQ131940–GQ132014. Of the 885 sites characterized, 768 were conserved across all 75 sequences and 82 of the 117 variable sites were parsimony informative. Under the HKY+I+$\Gamma$ model of sequence evolution, a transition (ti)/transversion (tv) ratio of 14.5 and base frequencies of $A = 0.3069$, $C = 0.1590$, $G = 0.1497$ and $T = 0.3844$ (% AT = 70.2%) were recovered. Third base position substitutions accounted for 88.03% of the variation with 11.11% and 0.85% being due to first and second base substitutions, respectively. Mutations at the nucleotide level gave rise to five non-synonymous amino acid substitutions at codons 4, 191, 219, 238, and 289. Of the 73 $B. huntleyi$ individuals sequenced, 62 had unique haplotypes corresponding to a haplotype diversity ($h$) of 0.995 and a nucleotide diversity ($\pi$) of 0.01278 for the Prince Edward Islands archipelago. When considering each island individually, 43 unique haplotypes were recovered from the 51 Marion Island specimens sequenced ($h = 0.992$ and $\pi = 0.01130$), whilst for Prince Edward Island, 19 unique haplotypes were identified from 22 individuals ($h = 0.987$ and $\pi = 0.00890$).

MP analysis recovered 76125 equally parsimonious trees with homoplasy indices of: $CI = 0.520$, $RI = 0.827$ and $RCI = 0.430$ when characters were assigned equal weights. Successive weighting with the $RCI$ recovered the same number of equally parsimonious trees and homoplasy indices of $CI = 0.699$, $RI = 0.889$ and $RCI = 0.621$. All methods of phylogenetic inference (Fig. 3) revealed the presence of two
distinct clades. Clade 1 (99–100% support) comprised three individuals from Marion Island collected from 1000 m a.s.l. that were basal to the second and larger clade comprising individuals from Marion Island and Prince Edward Island, suggesting that _B. huntleyi_ originally colonized Marion Island at high altitude. For the remaining individuals that grouped within clade 2, two major lineages could be discerned albeit with low levels of support. The first lineage (labelled ‘a’ in Fig. 3) contains individuals collected from Prince Edward Island alone, whilst lineage ‘b’ consists solely of individuals collected from Marion Island, the only exception being individual 317-19. Clade 2 is characterized by low bootstrap support for the internal nodes and by shallow divergences, with high support values occurring primarily at the terminal nodes. The haplotype network (Fig. 4) indicates that clade 1, the high altitude, relict population present on Marion Island, is separated from clade 2 by more than 12 steps. The two major lineages within clade 2, namely ‘a’ and ‘b’, which correspond to Prince Edward Island and Marion Island, respectively are separated from each other by more than seven steps.

Tests for rate heterogeneity revealed that _B. huntleyi_ lineages from the Prince Edward Islands archipelago do not

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**Fig. 5.** Mismatch distribution for _Bothrometopus huntleyi_ individuals from a. Marion Island (MI), and b. Prince Edward Island (PEI). Circles represent the simulated stepwise expansion model and the squares represent the observed data.

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**Table II.** Mismatch parameters for _Bothrometopus huntleyi_ from Marion Island (MI) and Prince Edward Island (PEI):

<table>
<thead>
<tr>
<th>Population</th>
<th>Mean number</th>
<th>Mismatch observed variance</th>
<th>τ (alpha = 0.100)</th>
<th>θ₀</th>
<th>θ₁</th>
<th>SSD</th>
<th>SSD Obs. (Sim. Ssd)</th>
<th>Raggedness index</th>
<th>P (Sim. Ssd)</th>
<th>P (Obs. Ssd)</th>
<th>θ₀/M</th>
<th>θ₁/M</th>
<th>P (Obs. Rag)</th>
<th>θ₀/M</th>
<th>θ₁/M</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEI</td>
<td>8.076</td>
<td>15.123</td>
<td>9.491 (5.764–12.577)</td>
<td>0.059</td>
<td>28.268</td>
<td>0.006</td>
<td>0.753</td>
<td>0.013</td>
<td>0.574</td>
<td>192478 (145282–337043)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MI</td>
<td>10.074</td>
<td>21.551</td>
<td>8.854 (6.683–15.594)</td>
<td>2.245</td>
<td>48.137</td>
<td>0.004</td>
<td>0.497</td>
<td>0.007</td>
<td>0.574</td>
<td>192478 (145282–337043)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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* Tests for rate heterogeneity revealed that _B. huntleyi_ lineages from the Prince Edward Islands archipelago do not
evolve at significantly different rates. The estimated time to
coealscence of *B. huntleyi* is c. 0.723 m.y.a., predating the
oldest dated rocks, but possibly not island emergence. The
first major inter-island dispersal event is estimated to have
occurred c. 0.216 million years (m.y.) later, at which time
the islands were fully formed, following which *B. huntleyi*
dispersed to the remainder of Marion Island during
predominantly glaciations-free periods. The results indicate
that subsequent to this range expansion, the populations on the
two islands have remained largely isolated with the only
evidence of inter-island gene flow being detected for a single
haplotype (317-19). This individual, sampled from Prince
Edward Island shares recent common ancestry with individual
355-1 from Long Ridge South (Fig. 3) on Marion Island, and
an estimated time to coalescence of c. 0.02 m.y.a. (Fig. 3).

The mismatch distributions for individuals collected on
each island are presented in Fig. 5. Neither the SSD nor the
raggedness index values (Table II) showed any significant
deviation from the expansion model, indicating a population
expansion for both Marion Island and Prince Edward Island.
The time of the main expansion events calculated using
\( \tau = 2 \mu t \) (Table II) corresponds to c. 0.2 m.y. before present (bp).

Fu’s *F*<sub>s</sub> statistic (Table III) showed statistically significant
and large negative values for both islands that are indicative
of a population expansion (Fu 1997). The *R*<sub>2</sub> statistic
(Table III) of Ramos-Onsins & Rozas (2002) also indicates
population expansion. The different estimations for \( \theta \) are
presented in Table III.

**Discussion**

Time to *B. huntleyi* haplotype coalescence on the Prince
Edward Islands archipelago is estimated to have occurred
3.703 m.y.a. This date is older than that of the oldest
dated rocks (0.45 m.y.a.), but remains within the estimated
emergence time of the islands of less than 1 m.y.a. (McDougall
*et al.* 2001, Boelhouwers *et al.* 2008). It is probable that the
founder population was restricted to high elevation areas of
Marion Island for a period of c. 0.216 m.y. High elevation
refugia in the form of nunataks have been identified for Marion
Island both on geomorphological and molecular grounds
(Boelhouwers *et al.* 2008, Mortimer *et al.* 2010). Moreover,
recent studies from Antarctica are also providing evidence of
refugia for invertebrate taxa that initially appear to be at odds
with the geological evidence (Convey *et al.* 2008).

*Bothrometopus huntleyi* is estimated to have then
dispersed to the remainder of Marion Island and on to
Prince Edward Island, approximately 0.507 m.y.a. This
occurrence coincides with the oldest dated rocks on the
island (Boelhouwers *et al.* 2008). Individual 317-19
from Prince Edward Island groups with individuals
(92% bootstrap support; Fig. 3) in the Marion Island-
associated lineage b in clade 2, and represents a lineage that
became established on Prince Edward Island much later at
c. 0.02 m.y.a. When comparing the dates estimated in Fig. 3
with the estimated times of glaciations and volcanic activity
on Marion Island (Fig. 1) it can be seen that, while numerous
divergence events coincide with times of volcanic activity,
very few divergence events occurred during the first and
third glaciations periods. Thus, it appears that genetic
divergence of *B. huntleyi* is linked to volcanic activity on
Marion Island, especially at those times when the island was
not glaciated. Quite what mechanism of transport was
involved is difficult to determine, but complex associations
between volcanism and population structure are well known
for animals from other island systems (e.g. Vandergragt

It should be noted that for *B. huntleyi* only one instance
of migration from Marion Island to Prince Edward Island
occurred after Prince Edward Island was originally colonized
(Fig. 1). This contrasts markedly with the multiple migration
events noted for *Ectemnorhinus* weevils from Marion Island
to Prince Edward Island (Grobler *et al.* 2006). The difference
between *B. huntleyi* and *Ectemnorhinus* weevils could be
explained by habitat preference. *Ectemnorhinus* weevils occur
mainly on vegetation (Chown 1989, 1992), and are thus
more likely to be transported by birds than *B. huntleyi* which
mainly occurs on rock surfaces and hides in crevices in the
rocks. *Bothrometopus huntleyi* individuals also have adhesive
brushes on the tarsomeres 1–3 (Kuschel & Chown 1995) that
prevent the weevils from being easily lifted by the frequent,
strong south-westerly winds.

The results of the mismatch distribution analyses are
consistent with that of the phylogeny as they indicate that
the major expansion event occurred on Marion Island
3.192 m.y.a., with lower and upper bounds of \( \pm 0.145 \)
to \( \pm 0.337 \) m.y.a., respectively. Most divergence times
calculated for Marion Island (Fig. 3) fall within the upper
bound estimates of the mismatch analyses. Similarly, most
Prince Edward Island divergence estimates (Fig. 3) also
fall within the lower and upper bounds of \( \pm 0.125 \) to
\( \pm 0.273 \) m.y.a., respectively as estimated by the mismatch
distribution analyses which revealed that a major expansion
event occurred on Prince Edward Island c. 0.206 m.y.a.
At present, the high elevation Marion Island founder population is intact (clade 1) and two island-discrete population complexes occur which are separated by an ocean barrier of 19 km. Historical inter-island migration has been minimal and unidirectional (from Marion Island to Prince Edward Island only) compared to other endemic Ectemnorhinus weevils studied to date. The isolation of the population on Prince Edward Island from the population on Marion Island probably resulted from a combination of factors, including a lack of both volcanic activity and of bird and/or wind-assisted dispersal between the islands. The genetic data are thus consistent with the lack of morphological differences between the individuals found on the islands, and the assumption of a shared species. Whilst the phylogeny did not recover highly-supported island-specific clades, the haplotype network (Fig. 3) revealed that B. huntleyi individuals from Prince Edward Island are genetically more similar to each other than to individuals from Marion Island. It is therefore likely that these allopatric populations will continue to diverge as long as the natural inter-island isolation is maintained.

An important conservation biogeographic insight from our work is that anthropogenic gene flow between populations on Marion Island and Prince Edward Island should be prevented. Thus, the current practice of limiting visits to Prince Edward Island to a minimum, and the strict quarantine provisions for such visits (e.g. Davies et al. 2007) should be maintained. Moreover, the genetic distinctiveness of the B. huntleyi population on Marion Island, which is under threat via increased size-selective predation by the house mouse, Mus musculus domesticus (Chown & Smith 1993), also underscores the need to control mice on Marion Island to ensure that both the unique genetic variation on Marion Island, and the evolutionary dynamics of the archipelago are conserved.

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