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1. List of Broscini species used in this study with information regarding authority, distribution (S.I. = South Island New Zealand, N.I. = North Island New Zealand; Ch.Is. = Chatham Islands) and number of individuals employed.

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(M.= MecodemaMeta= Metaglymma O.= Oregus D.= Diglymma S.I. = South Island New Zealand; N.I. = North Island New Zealand; BOP = Bay of Plenty; genes analysed: a = COI, COII, 16S, 18S; b = COI, COII, 16S; c = COI, 18S; d = COI; LUNZ = Lincoln University Entomological Research Museum; P = S.M. Pawson collection); COI sequences from samples of the ecodemaingroup marked with * have previously been deposited in Genbank#

naturally an increasing interest in how diversification is distributedare predatoryMecodenaa diverse genus with species distributed through time and space. throughout the New Zealand mainland from alpine to coastal

In this study we examine the phylogenetic relationships and abitats. In contrast, there is a single speolexe odema alternams timing of radiation ofMecoden blanchard, 1843) carabid beetles the Chatham Islands. The same species occurs in southeast New (tribe Broscini). This endemic genus of large, flightless beetlesealand near Dunedin. AlthoughM. alternansnay be better constitutes a prominent species radiation in New Zealand andreated as a species complex [47], no morphological characters presents a good opportunity to explore species level diversificatiohave yet been described that distinguish Chatham Island We utilise the fact that the genus is represented on the Chathamopulations from those in mainland New Zealand ([49] & I. Islands, which are located approximately 850 km east of the South Townsend pers. obs.).

Island, New Zealand in the Pacific Ocean. Geological evidence for In total our sampling comprised 113 specimens, with 88 the formation of this archipelago within the last 4 Myr is Mecodemapresenting 35 described species, and 4 undescribed compelling [3,32,33,34] and corroborated by genetic data forspecies of the 66 recognizedbecodemagecies (after [45,46] and many taxa (e.g. insects – [18,35], plants – [36,37,38], parakeets http://www.landcareresearch.co.nz/research/biosystematics/ [39], pigeons – [40,41], cicadas – [42], invertebrates and plants – invertebrates/carabid/carabidlist), see Table 1 for details and [43]). In this study the earliest possible establishment of an islanauthorities. Putative outgroup New Zealand Broscini in our biota (4 Myr) on the Chathams [32] is used as a maximumsample includedOregutPutzeys, 1868DiglymmaSharp, 1886), possible calibration for estimating the timing of diversification inBrullea antarcti(Laporte de Castelnau, 1867), etaglymm(Bates, Mecodema urthermore a substitution rate for Coleoptera [44] is

employed to further explore timing of lineage diversification of this beetle genus.

Materials and Methods

Sampling

Table 2.

The genusMecodem(Blanchard, 1853) belongs to the tribe Broscini (Carabidae). Broscini has a worldwide distribution but has its main diversity in the southern hemisphere (subfamily Nothobroscinae) [45] and consists of at least 27 genera, comprising about 80 species (see http://www.landcareresearch.co.nz/research/ biosystematics/invertebrates/carabid/carabidlist) [46,47]. Six endemic genera of Broscini are recognized in New Zealand, but Mecodema especially species-rich. Adultecodemabetles are relatively slow-moving, nocturnal, flightless (with fused elytra), generally active throughout the year, and usually scarce [48]. As with other Carabidae, adults and larvae of the New Zealand taxa

1867), and Bountya insula ris wnsend, 1971); plus one representative of Broscini from AustraliaChylnus at (Putzeys, 1868) (Table 2). As many of the species Niecodemare scarce and therefore difficult to collect we also made use of material from museum collections, and supplemented outgroup sampling with available GenBank sequences (gus septentrionalis66847 & AF466848; Oregus crypticulates 4423; Oregus inaequalis 466850 [50]; Calathus aztec U254333 [51]; Broscosoma relictem 2502; Promecodesps AF012499Creobius eydoux012498 [52]). The resulting sample represents the geographic and ecological range of Mecodemia New Zealand (Table 2)Mecodemspecies show examples of allopatric, parapatric and sympatric distribution. Species but not always species groups are limited to particular areas and only few species (e.g. crenicolle

k incellihood phylogensy for M c **a.** A) Spatial distribution of samples used in this analysis. Symbols correspond to those in Fig. 2B and code for different clades. B) Analysis of concatenated dataset including mitochondrial COI, COII and 16S plus nuclear 18S. Values at nodes indicate ML bootstrap support returned by analysis using RaxML. Specimen numbers at tips are given as in Table 2. doi:10.1371/journal.pone.0086185.g002

Polymerase chain reactions were performed im 10 0 lumes Phylogenetic analysis

and the amplified products then checked on a 1% agarose gel and To test whetherDiglymmaand Oreguspecies are the natural purified using SAP/EXO1 digest (USB Corporation). Purified sister group toMecodemae first analysed data from two genes PCR products were sequenced using standard protocols for the eparately (COI and 18S) as it was not possible to gain sequences ABI Prism BigDye Terminator Ready Reaction Kit (Applied for outgroup taxa outside of New Zealand for all the employed Biosystems, Mulgrave, Australia) and run on an ABI Prism 377_{Species} and sequence availability in GenBank within Broscini was automated sequencer (Applied Biosystems). Sequence identity waso very poor. AlthougbreguandDiglymmæpresent two of the confirmed by comparison with published data, checked for New Zealand Nothobroscina genera considered closest to nucleotide ambiguities in Sequencher 4.2 (Gene Codes Corpora_{decoden} [45], it was crucial to verify this relationship within tion, Ann Arbor, MI, www.genecodes.com) and aligned using SeBroscini as two other potential outgroup taxaletaglymmand AI v2.0a11 [58]. The sequences have been deposited with enlige exist. MrBayes 3.1.2 [59] was used to implement Bayesian accession numbers KF913050–913193 at GenBank (16Sanalysis with the datasets applying a GTR model with gamma-KF913050–913088; 18S: KF913089–913130; COII: KF913131– distributed rate variation across sites and a proportion of 913169; COI: KF913170–913193). invariable sites. Analyses with MrBayes used four independent

 $f M c a$ $f t A$) COI Bayesian tree generated in BEAST. Numbers on nodes show age estimates based on stratigraphic calibration of 4 Myr. Outgroup taxa were the same as used in the outgroup test (Fig. 1). Grey branches indicate lineages present in North Island New Zealand, and black branches indicate South Island lineages. Tip symbols correspond to clade and location identifiers in Fig. 2. Coloured symbols match symbols in Fig. 3B, highlighting disjunct lineages in southern North Island and northern South Island. Asterisks indicate ag e estimates between N.I. and S.I. lineages (*=1.60 Myr, **=1.69 Myr, ***=2.03 Myr). B) Reconstruction of the paleogeographic environment in lower North Island, New Zealand ca. 3 million years ago, green areas indicating likely land above sea level during this time (modified from [28]). Black outlines indicate present day New Zealand land area with coloured symbols corresponding to those in Fig. 3A, showing the present sampling

4. BEAST time estimates based on stratigraphic and COI substitution rate of Coleoptera [44] calibration.

	(5%) ttt \mathbf{t} t	(5%) tt t t t t
A	4	0.2 $(0.0 0.46)$
B	$1.24(0.5 - 2.09)$	$0.07(0.02 - 0.15)$
C	$8.38(5.29 - 12.06)$	$0.52(0.22 - 0.9)$
D	$6.39(3.97 - 9.41)$	$0.4(0.18 - 0.72)$

Markov Chain Monte Carlo (MCMC) runs for ten million alternansnd its closest relative on mainland New Zealand. We generations with a burn-in of 10% and a tree sampling frequencyassumed a normal distribution for the age around a calibration of 1000. Results were checked for convergence. Resultingalue of 4 Myr, derived from the maximum age for the Chatham posterior probabilities on the nodes were recorded. Islands land surface [3,33], assuming that colonization was most

To examine the species phylogeny of the codematoup in New Zealand we employed all four genes (three mitochondrial an做vely, to capture the minimum likely diversification dates we also one nuclear) with a subset of 50 taxa (44 ingroup and 6 outgroupalibrated the COI dataset with the substitution rate estimated by samples). The outgroup sampling was chosen after considerationons et al. [44] for Coleoptera COI. This included a normally of the results from the prior outgroup analyses. All taxa with datadistributed prior on the substitution rate of 0.08606 subst/site/ missing for no more than one of four genes were included in thenyrs/l, and a 95% HPD interval from 0.0253–0.147 subst/site/ phylogenetic analysis (Table 2). Partition-homogeneity tests (PHT myrs/l as an approximation to the posterior distribution provided [60]) were implemented in PAUP*4.0b10 [61] with 500 replicates by Pons et al. [44]. This rate obtained from analysis of numerous for the combination of the gene regions to detect significan peetle taxa is amongst the highest estimated for any animal gene, heterogeneity among the data sets. likely sooner after emergence of the islands than later. Alternaand other rates obtained for particular beetle lineages are much

MrBayes 3.1.2 [59] was then used to implement Bayesian^{slower} (e.g. 0.0211 subst/site/myrs/l [64], 0.02 subst/site/myrs/l analysis with the concatenated dataset, applying a GTR mode^[65]). Even these rates are nearly twice the widely employed with gamma-distributed rate variation across sites and a propor^{0.0115} subst/site/myrs/l estimate of Brower [66]. Age estimation tion of invariable sites. The same model was applied to the or both datasets and both calibration strategies were conducted partitions with rates and nucleotide frequencies for each gene^{nder} the assumption of a strict molecular clock as well as unlinked. Analyses with MrBayes used four independent Markov^{assuming} a relaxed molecular clock with a lognormal distribution Chain Monte Carlo (MCMC) runs for two million generations of rates along the phylogenies [67]. The fit of both priors was with a burn-in of 25% and a tree sampling frequency of 1000. compared using Bayes Factors. For all datasets and calibration Resulting posterior probabilities on the nodes were recorded. The trategies the relaxed lognormal clock fitted the data decisively same data were subjected to Maximum Likelihood analysis with the than the strict clock [68].

bootstrap resampling incorporating a GTR model with gammapartitioned by gene (COI, COII, 16S, 18S) and bootstrap starting tree under the GTRI+C model of nucleotide substitution. resampling was halted by RaxML

Molecular dating

As fossil remains *di*lecodentizat could provide information for for Coleoptera COI [44] for calibration. In order to gauge the timing and extent of species radiation Mecodemaithin New Infinity and extern or species radiation intecodemation new MCMC was confirmed, the posterior distributions of all param-
Zealand COI sequences were obtained for additional specimens intersection are estimated from the com addition to previous analyses (Table 2). In some cases, this dreal runs conducted for each analysis. The program FigTree 1.4.0 upon museum specimens to further assess the stability of orthogo was used to visualize the reconstructed phylogenies. inferences about the distribution of diversity and timing of radiation in this beetle group.

distributed rate variation. ML analysis used RaxML [62] [70] for molecular dating with the given calibrations. All analyses implemented via the CIPRES portal [63]. The data were were conducted with a Birth-Death tree prior and a random We used the software BEAUTI 1.4.8 [69] and BEAST 1.7.5 The MCMC was run for 50–100 million generations, sampling

As lossilied and subcodernal codid provide information for distributions of all parameters in question, to verify convergence
Calibrating a molecular clock have not been found, yet, we had to the MCMC and to estimate Effec rely on geological information and a substitution rate calculated in the mome and to cumulate Encentre Cample Size (ECC). In the every 5000-10,000th step after a discarded burn-in of 5-10 million steps. Each analysis was run at least two times. The program Tracer 1.4 [71] was used to summarize posterior of the MCMC and to estimate Effective Sample Sizes (ESS). If the conducted for the respective analysis. After convergence of the eters were estimated from the combined posterior distributions of

In total 113 unique COI sequences were used for molecular dating in Mecodem_{(Table 2}). To obtain estimates for the maximum age of lineage formation within the genus codema to gauge the scale of genetic diversity among lume codema we used this dataset of COI with two different calibration specimens. Overall we observed relatively low genetic distances strategies. First we employed a stratigraphic calibration to estimatamong species dMecodemaith a maximum ML-distance of divergence times using the split between Chatham Island 0.0179 in COII (COI: 0.0161, 16S: 0.053). Lower values for 16S \mathbf{t} Three widely used mitochondrial gene regions were employed

compared to COI and COII reflect the comparatively low proportion of variable sites in this gene (16.9%).

Separate analyses of COI and 18S DNA sequences from 41 specimens ofMecodemaand 6 outgroup taxa (Table 2) resulted in similar topologies even though sequence variation in 18S was low. These analyses confirm@reguand Diglymmas the sister group to Mecodemand revealed the placement Metaglymmaad Brullea within the Mecodennadiation (Figs. $1A & B$).

The alignment of data from four gene regions comprising 50 specimens sampled across New Zealand (including 6 outgroup specimens –Table 2, Fig. 2) was 3114 bp long in total. All three mitochondrial genes displayed the average insect A-T content of about 75%. The partition homogeneity test (PHT) revealed no significant heterogeneity of lineage partitioning among the data sets $p = 0.866$, suggesting their concatenation was appropriate. The GTR+I+C model of nucleotide substitution was identified as the best fitting model by the hLRT and the AIC as implemented in Modeltest 3.5 [72].

Bayesian and ML analysis of the concatenated dataset supported a single topology with the same groupings and branching order and well-supported nodes (FigMa)codemass confirmed as paraphyletic with respect to that and Brulleas these fall inside the Mecodemamplex throughout all datasets and analyses in this study. We note that in all cases aglymmand Brulleaare placed within theM. curvidens/origoides up. This phylogenetic position contradicts the current taxonomic classification and needs to be addressed further in the future. Clades revealed in this analysis comprise species that are, in many cases, by the setting of priors. In this study the inferred mitochondrial divergence times based on the fast COI rate obtained for Coleoptera [44] are consistently substantially younger than would be expected for a highly diversified genus even in the relatively youthful landscape of New Zealand.

Despite a perception that New Zealand is an ancient land mass

demonstrating the population genetic and ecological mechanismatergenerational population genetics of large invertebrate popuof diversification (e.g. [89]). lations will provide the basis of exciting research.

Increasingly, the fields of species phylogenetics and population phylogeography are merging as it becomes easier to generate $\mathbf k$ appropriate DNA data, and the focus in taxonomy is shifted towards an evolutionary paradigm (e.g. [90]. Teasing apart the \mathbb{R} k t We would like to thank Steve Pawson and Peter Johns for contributing

Interaction of abiotic and genetic processes on population plushes are assigned to this study. Mary Morgan-Richards and Frank Wieland gave subdivision remains challenging but codenina one taxon group that will provide helpful insight, and it is already evident that Conservation assisted with permitting. valuable comments on earlier drafts of the manuscript. The Department of

Mecodema an impressive example of recent species radiation in the New Zealand fauna. In recent years, synthesis of phylo^{A} **t t t**

genetic, ecological and taxonomic evidence has indicated thatonceived and designed the experiments: JG SAT. Performed the the biology of New Zealand is primarily the story of recent experiments: JG. Analyzed the data: JG MK SAT. Contributed adaptation and speciation [8,9,19]. Understanding properly reagents/materials/analysis tools: JG MK RME JIT SAT. Wrote the how our view of "recent" geological time relates to the paper: JG SAT.

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