# Subtle limits to connectivity revealed by outlier loci within two divergent metapopulations of the deep- sea hydrothermal gastropod *Ifremeria nautilei*

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### Abstract :

Hydrothermal vents form archipelagos of ephemeral deep-sea habitats that raise interesting questions about the evolution and dynamics of the associated endemic fauna, constantly subject to extinction-recolonization processes. These metal-rich environments are coveted for the mineral resources they harbor, thus raising recent conservation concerns. The evolutionary fate and demographic resilience of hydrothermal species strongly depend on the degree of connectivity among and within their fragmented metapopulations. In the deep sea, however, assessing connectivity is difficult and usually requires indirect genetic approaches. Improved detection of fine-scale genetic connectivity is now possible based on genome-wide screening for genetic differentiation.

Here, we explored population connectivity in the hydrothermal vent snail Ifremeria nautilei across its species range encompassing five distinct back-arc basins in the Southwest Pacific. The global analysis, based on 10 570 single nucleotide polymorphism (SNP) markers derived from double digest restriction-site associated DNA sequencing (ddRAD-seq), depicted two semi-isolated and homogeneous genetic clusters. Demo-genetic modeling suggests that these two groups began to diverge about 70 000 generations ago, but continue to exhibit weak and slightly asymmetrical gene flow. Furthermore, a careful analysis of outlier loci showed subtle limitations to connectivity between neighboring basins within both groups. This finding indicates that migration is not strong enough to totally counterbalance drift or local selection, hence questioning the potential for demographic resilience at this latter geographical scale. These results illustrate the potential of large genomic datasets to understand fine-scale connectivity patterns in hydrothermal vents and the deep sea.

**Keywords** : ddRAD-seq, Demographic inference, Genetic connectivity, Hydrothermal vents, Outlier detection, Western Pacific

# Introduction

Understanding the connectivity of populations is a central issue for evolutionary ecology, conservation and management (Cayuela et al., 2018). Direct approaches such as population monitoring or mark-recapture experiments are rarely applicable in marine environments, because many marine species have large population sizes and high dispersal capabilities due to their minute pelagic propagules. These characteristics are likely to reduce the ability of population genetics to assess population connectivity at local and regional scales, except in situations where there is sufficient genetic differentiation or where a large fraction of the population can be sampled (Jones et al., 2005; Pinsky et al., 2010). Deep-sea hydrothermal ecosystems have attracted much attention since their discovery in the late 1970s (Van Dover et al., 2001; Tunnicliffe et al., 1998) due to the oasis-like distribution of these unique and chaotic habitats harboring rich and endemic fauna. Hydrothermal environments are mostly found in tectonically active areas, such as mid-oceanic ridges, where neighboring vents are often separated by tens of meters to hundreds of kilometers, resulting in an almost linear, but fragmented and unstable distribution of vent communities (Chevaldonné et al., 1997; Hannington et al., 2011; Jollivet et al., 1999; Vrijenhoek, 2010).

Studies of slow (*e.g.* Mid-Atlantic Ridge) and fast (*e.g.* East Pacific Rise) spreading ridges have shown that number of species are able to maintain gene flow over thousands of kilometers (Breusing et al., 2016; Craddock et al., 1995; Hurtado et al., 2004; Jollivet et al., 1995; Teixeira et al., 2011, 2012; Won et al., 2003; Yahagi et al., 2019). Many vent invertebrates possess longrange planktonic larvae that can rapidly (re)colonize newly formed sites (Mullineaux et al., 2010). This high larval dispersal capacity leads to local colonization processes following a stepping-stone mechanism of exchanges, or to the formation of more complex metapopulation dynamics where local extinctions and migration may vary greatly according to the geotectonic context of the venting sites (Audzijonyte & Vrijenhoek, 2010; Jollivet et al., 1999; Vrijenhoek, 1997, 2010). Population connectivity can, however, be severely hampered by physical barriers to larval

dispersal such as transform faults, diverging ocean currents or microplates (Johnson et al., 2006; Plouviez et al., 2009; Plouviez, Schultz, et al., 2013).

The hydrothermal ecosystems found in the Southwest Pacific are mainly associated with the formation of back-arc basins (BABs). BABs result from complex subduction processes between several plates, leading to a discontinuous and nonlinear distribution of venting sites. Hence, the question arises as to the degree of connectivity between populations inhabiting these BABs, noticeably to address the issue of their resilience with respect to deep-sea mining projects (Gena, 2013; Niner et al., 2018). In this context, only a few studies to date have focused on understanding the general patterns of spatial genetic connectivity in ecologically vulnerable hydrothermal species. For instance, Thaler et al. (2011) showed that the gastropod Ifremeria nautilei is genetically differentiated between the Manus and North-Fiji/Lau basins. Similar results have been reported in other species, such as the limpet Lepetodrilus schrolli (Plouviez et al., 2019), the shrimp Rimicaris variabilis and the squat lobster Munidopsis lauensis (Thaler et al., 2014). Moreover, this latter species is characterized by additional intra-basin structuring. In contrast, the limpet Shinkailepas tollmani does not show any differentiation at any of these scales (Yahagi et al., 2020). However, due to the use of only a limited number of markers, none of these studies have reached the resolution necessary for the fine-scale assessment of connectivity in these species.

During the last decade, the development of next-generation sequencing (NGS) and associated techniques have increased the quantity and accessibility of population genomic data, particularly in non-model species. Analyzing these large datasets with thousands of markers along the entire genome using demo-genetic inference methods helps reveal the complex demographic histories of species (Excoffier et al., 2013; Feutry et al., 2020; Gutenkunst et al., 2009; Rougeux et al., 2017; Tine et al., 2014). NGS datasets also give access to unprecedented statistical power to detect non-neutral genetic variation (outlier loci) that can potentially provide finer scale spatial information on connectivity, dispersal and possibly local adaptation (Gagnaire et al., 2015; Milano et al., 2014; Wyngaarden et al., 2017). This information can potentially help distinguish situations of genetic connectivity—whereby local populations are demographically independent (*i.e.* mainly replenished by local propagules) but long-range gene flow mediated by a small number of propagules is sufficient to ensure the circulation of genetic variation among them—from situations of demographic connectivity where a substantial fraction of local population size is made up of

immigrants (Lowe & Allendorf, 2010). The consequences of this difference in population connectivity in terms of resilience to local extinction or habitat destruction are quite obvious, with prompt recolonization being expected only in the second situation.

The aim of the present study was to elucidate fine-scale population structure and connectivity using high-throughput double-digest restriction-site associated DNA (ddRAD) sequencing on Ifremeria nautilei, whose known distribution covers the Southwest Pacific, from the Manus BAB in Papua New Guinea to the Lau BAB off the Tonga Islands. This hydrothermal gastropod of family Provannidae harbors chemoautotrophic symbiotic bacteria in its gills to produce organic matter and forms dense aggregations around diffuse fluid venting at temperatures lower than 15°C (Borowski et al., 2002; Windoffer & Giere, 1997). The species is gonochoric with a nearly equal sex ratio and females brood their progeny in a metapodial pouch until the lecithotrophic embryos (several thousands of similar size) reach a specific and unique pre-veliger stage known as Waren's larva (Reynolds et al., 2010; Warèn & Bouchet, 1993) . The gastropod reproduces via internal fertilization leading to a patchwork of brooding and nonbrooding females throughout the year due to asynchronous spawning. Brooding is a reproductive trait that usually limits the dispersal ability of species; however, the lifespan of *I. nautilei* pelagic larvae is not known, nor is the maximum distance veligers can travel prior to settlement on a new venting site. Constituting a large portion of the biomass and harboring other species such as S. tollmanni and L. schrolli, I. nautilei is a keystone species important for these deep-sea ecosystems. Furthermore, it is classified as endangered by the IUCN (https://www.iucnredlist.org/species/145380421/145380604) along with some other hydrothermal species. Therefore, studying the connectivity patterns of *I. nautilei* is a flagship endeavor to assess the potential impact of deep-sea mining on this keystone species and its associated fauna.

For this study, extensive sampling was carried out on 29 sites in 17 hydrothermal vent fields across five basins distributed over 5000 km in the Southwest Pacific.

# **Materials and Methods**

Sample collection and DNA extraction

A hierarchical sampling plan was deployed on board the French oceanographic vessel RV *L'Atalante* during the ChubacArc 2019 oceanographic cruise using the remotely operated underwater vehicle (ROV) *Victor 6000*. A total of 684 individuals were collected in the Southwestern Pacific Ocean from 29 sampling sites distributed among 17 vent localities or hydrothermal fields across four BABs and one volcanic submarine area (Futuna), spanning the entire known geographical distribution range of *I. nautilei* (Figure 1, SI Table S1). This sampling includes samples from the newly discovered active site La Scala in the Woodlark basin (Boulart et al., in press).

In addition, 27 individuals were added from collections of previous oceanographic cruises, with 22 samples obtained by S. Hourdez during the Lau basin 2009 oceanographic cruise with few at the now-extinct Kilo Moana site and 5 samples obtained during the Manus basin 2009 cruise kindly provided by C. L. Van Dover (SI Table 1). Altogether, a total of 456 unique samples were used and analyzed in this study, of which 362 remained after filtering the sequence dataset (SI Table 2).

Once on board, the snails were dissected and various tissues were preserved in EtOH or frozen at -80°C. Genomic DNA was extracted from fresh foot tissue to limit DNA contamination by symbionts hosted in the gills. A fraction of the tissue samples was stored in 90% EtOH for backup. Samples from C. L. Van Dover's collection were preserved in 90% EtOH, and those from S. Hourdez were kept frozen at -80°C. Genomic DNA was extracted using the NucleoSpin Tissue kit according to the manufacturer's protocol (Macherey-Nagel); some samples were extracted using a standard phenol-chloroform method.

### Preparation and sequencing of ddRAD libraries.

DNA extracts were visualized on 0.8% agarose gels and each concentration was standardized to between 10 and 50 ng. $\mu$ l<sup>-1</sup> after a fluorometric quantification with the QuantiFluor dsDNA system (Promega). Individual double-digest restriction-site associated DNA (ddRAD) *PstI-MseI* libraries were prepared following (Brelsford et al., 2016) after modifications detailed in (Daguin-Thiébaut et al., 2021). Five pooled libraries were prepared with a combination of four to eight Illumina indexes and 24 barcodes per index, multiplexing a total of 486 samples, including 27 pairs of replicates for quality control, representing 456 individuals. The sequencing effort was sized to produce on average  $3 \times 10^6$  read pairs per individual. Each genomic pool was sequenced on one

lane of a HiSeq4000 Illumina sequencer (paired end, 150 bp) at the Genoscope sequencing facility (Centre National de Séquençage, Evry, France).

### De novo ddRAD-tag assembly, SNP calling and filtering

Fastqc (V.0.11.9) was used only for quality control of raw reads, no filters were applied on them. Individual reads were demultiplexed using the "Processradtag" pipeline in Stacks (V.2.52) (Rochette et al., 2019). Due to the lack of a reference genome for *I. nautilei*, reads were assembled *de novo* using each Stacks module one by one (ustacks, cstacks, sstacks, tsv2bam, gstacks and populations). To identify the most appropriate assembly parameters, we followed previously published recommendations (Mastretta-Yanes et al., 2015; Paris et al., 2017) (See SI and SI Figures 1 to 6 for details). Briefly, we used the genotyping error rate between replicates, the number of variants (SNP), polymorphic loci (ddRAD-tags) and nucleotide diversity ( $\pi$  estimated in Stacks) as a function of the assembly parameters (m and M = n) determined with a subset of individuals covering all basins and localities (n = 84). The selected parameters were as follows: m = 4 (the minimum number of reads to assemble a stack), M = 8 (the maximum number of mismatches between putative alleles within individuals), n = 8 (the maximum number of mismatches between putative loci within the catalog of individuals) and R = 0.8 (the minimum percentage of individuals sharing a locus across all populations in the "populations" module).

After *de novo* assembly, several filters were applied using VCFtools (V.0.1.16) (Danecek et al., 2011) to reduce missing data and to account for potential paralogs (see SI Table 3). Briefly, we removed 1 of each of the 27 replicated individuals with the highest value of missing data. Then, we excluded SNPs with heterozygosity > 0.6, SNP and only individuals with less than 10% missing data were kept. Variants with a mean coverage higher than 80X were excluded. Using VCFtools, we excluded loci with a minor allele frequency (MAF) lower than 5% (alternative allele), followed by those that deviated significantly from Hardy-Weinberg equilibrium (*p*-value  $\leq$  0.05). We then kept only one randomly chosen SNP per ddRAD-tag to avoid short distance linkage disequilibrium between SNPs. PGDSpider (V.2.1.1.5) (Lischer & Excoffier, 2012) was used to convert the final VCF into the formats required for subsequent analyses.

### Population structure and diversity

Principal component analysis (PCA) was first performed on the final dataset with the R package SNPrelate (V.1.24.0) (Zheng et al., 2012). Pairwise fixation indices ( $F_{ST}$ ) were calculated in

Arlequin (V.3.5.2.2) (Excoffier & Lischer, 2010). AMOVA (Excoffier et al., 1992) was performed with 10 000 permutations of genotypes between populations by considering hierarchical geographic structure of localities within basins (See SI for parameters). Co-ancestry analyses were performed through ADMIXTURE (V.1.3.0) (Alexander & Lange, 2011) with 10 independent runs for K = 1 to 5. The best K value was selected by using the cross-validation error as recommended by the authors. Runs of ADMIXTURE were grouped using CLUMPAK (Kopelman et al., 2015), graphical visualizations of the results were plotted using library ggplot2 (V.3.3.3) in R (V.4.0.1). TreeMix (V.1.13) (Pickrell & Pritchard, 2012) was performed with 10 independent runs with migration events ranging from 0 to 5. The optimal number was selected according to the loglikelihood of each model.  $F_3$  admixture tests (Reich et al., 2009) were done using the THREEPOP programs implemented in TreeMix (V.1.13) package with default values.

To detect potential kinship, SNPrelate was used to compute identity-by-state between pairs of individuals. We used this approach to (1) minimize the risk of labeling error/exchange during the process of library construction and sequencing and (2) infer the level of kinship structure between non-replicated individuals because the existence of undetected underlying kinship structure can distort the population structure estimated by the gene genealogies.

Observed heterozygosity ( $H_o$ ), expected heterozygosity ( $H_e$ ), heterozygote deficiency ( $F_{IS}$ ), nucleotide diversity ( $\pi$ ) and raw nucleotide divergence ( $D_{xy}$ ) were calculated with the population module of Stacks using all sites from all ddRAD-tags in the final dataset.

### Demo-genetic history of the species

The demographic history of the species targeting past and present gene flow between metapopulation clusters was inferred using a modified version of  $\partial a \partial i$  (V.2.1) (Diffusion Approximations for Demographic Inference; Gutenkunst et al., 2009), with a dual annealing optimization function. This software simulates the joint allele frequency spectrum (JAFS) of two (or more) interacting populations according to different demo-genetic scenarios. Here, we considered 28 distinct scenarios built according to the population models used in Rougeux et al. (2017) with very few modifications, detailed below.

Basically, all these models derive from four basic models representing strict isolation (SI), isolation with migration (IM), ancient migration (AM), and secondary contact (SC). Each of them consists of an ancestral population of  $N_{anc}$  size that splits into two sister populations of effective

size  $N_1$  and  $N_2$  during time Ts for the (SI) and Tsm for the (IM) model, Tam+Ts for the (AM) model and Ts+Tsc for the (SC) model, where Ts is the time spent since the split of the two populations without migration, Tsm, the time spent since the split of the two populations with migration, Tam, the duration of the ancient migration period after the split of the ancestral population and before the emergence of strict isolation (Ts) and Tsc the duration of a secondary contact after a period Ts of strict isolation. Directional migration between populations is allowed at rates  $m_{12}$  and  $m_{21}$  from population 2 to population 1 and vice versa.

Further complexity was introduced as in Rougeux et al. (2017), by adding several processes occurring after the split, such as population expansion or contraction (G), the effect of linked selection reducing the effective population size of loci over a certain fraction of the genome (2N) and the effect of semipermeable genetic barriers (*i.e.* partial reproductive isolation) reducing the effective migration rate of loci over a certain fraction of the genome (2m). Furthermore, to dissociate the effect of the effective population size (genetic drift) and migration (gene flow), we only allowed the growth parameters (G) to vary during the migration phase of each model. Graphical representation of the four basic models and the three more complex models are displayed in SI Figure 7.

For the input dataset, we considered the two metapopulations defined by the global analyses (see Results), which corresponded to Manus/Woodlark and North-Fiji/Futuna/Lau. We used the folded joint allele frequency spectrum (folded JAFS), because no external group was available to identify the allelic ancestral states. All models were fitted independently of the dataset using dual-annealing optimization and run 10 times independently each to check convergence. Model comparisons were based on the Akaike information criterion (AIC). Using the best selected models, we then converted demographic parameters into biological units. In the absence of precise information on mutation rate and generation time for this species, we used  $10^{-8}$  as the mutation rate per site per generation. This widely used value falls within the range proposed by (Lynch, 2010), although admittedly the real value may be much larger or much smaller, as recalled in the Discussion. Parameters estimated using  $\partial a \partial i$  are scaled from the ancestral effective population size (N<sub>anc</sub>), which was estimated using the following formula:

$$N^{anc} = \frac{\theta}{(4 \times \mu \times L)}$$

where *L* represents the total length of the DNA sequence used in  $\partial a \partial i$ :

$$L = \frac{z \times y \times 275}{x}$$

where z represents the number of SNPs used, y the number of RAD-tags of 275 bp, and x the initial number of SNPs (z = 17365, y = 17365 and x = 250502, L = 331032).

Estimated times were calculated in units of  $2*N_{anc}$  generations and the migration parameters (m<sub>12</sub> and m<sub>21</sub>) were divided by  $2*N_{anc}$  to obtain the number of migrants in each population per generation. The standard deviations were estimated using the Fisher information matrix (FIM) method implemented in  $\partial a \partial i$ .

### Outlier loci and detection of fine-scale structure

To test whether fine-scale genetic structure exists within each genetic cluster defined as a result of the global analyses described in the preceding paragraph (see Results for details), we used several genome-scanning methods to identify candidate outlier SNPs (*i.e.* loci showing higher or lower levels of differentiation than expected under assumed neutrality). Such loci may be informative about fine-scale population structure and connectivity patterns (Gagnaire et al., 2015). Four different outlier detection approaches were used. The rationale behind this multiple testing is that these methods operate with somewhat different underlying assumptions or test statistics and are known to have varying discriminatory power depending on the situations to which they are applied (Villemereuil et al., 2014). Outlier loci were selected according to statistical thresholds (*p*-value  $\leq$  0.05 and 0.01) in each software package, while checking that candidate loci outnumbered the number of loci expected to fall outside the distribution by chance only (false positives). Furthermore, to focus on the relevant scale and avoid the detection of false positives due strong geographic structuring, these programs were run independently on each Manus/Woodlark and North-Fiji/Futuna/Lau metapopulation previously defined in the global analyses, while considering populations either by basin or by locality within these groups.

Four methods were used. (1) BayeScan (V.2.0) (Foll & Gaggiotti, 2008) detects potential outlier loci by using differences in allele frequencies under a simple island model in a Bayesian framework. Five independent runs were performed with the default settings. (2) PCAdapt (V.4.3.3) (Luu et al., 2017) uses the correlation of SNPs with the first principal components of the PCA to detect outliers by computing a Mahalanobis distance between their z-score on each PC. (3) Arlequin (V.3.5.2.2) (Excoffier & Lischer, 2010) detects outlier SNPs under a non-hierarchical

finite island model integrating  $F_{ST}$  and heterozygosity through 20 000 coalescence simulations of the neutral distribution with 100 demes each. (4) The core model of Baypass (V.2.1) (Gautier, 2015) based on a hierarchical Bayesian model in which loci that are more differentiated than expected under a non-equilibrium drift model are identified through the distribution of a statistic similar to  $F_{ST}$  corrected to account for demographic history. Baypass was run five times independently with default settings under the core model.

PCA, ADMIXTURE and  $F_3$  tests were then performed on the various outlier subsets to explore the information they convey.

In addition, outlier loci identified at the threshold ( $p \le 0.05$ ) were first blasted (BLASTN, E-value threshold: 10<sup>-5</sup>) against the *Alviniconcha boucheti* transcriptome, which was previously assembled using rnaSPAdes (V.3.13.1) (Bankevich et al., 2012) (cf. Castel et al., in prep). Transcript hits with a size greater than 300 bp were subsequently blasted (BLASTX, E-value threshold: 10<sup>-5</sup>) against the NCBI UniProtKB/Swiss-Prot database using the software Geneious Prime® 2021.2.2.

### Results

### De novo assembly and data filtering

*De novo* assembly resulted in a dataset of 38 608 ddRAD-tags with a mean coverage of 14X for 486 samples. The mean genotyping error rate was 0.48% and the maximum value was 1.06% from all pairs of replicates. These ddRAD-tags contained 649 106 SNPs. Following SNP filtering, the final dataset resulted in a VCF file containing 362 individuals with 10 570 unlinked bi-allelic variants with an individual mean coverage of 17.7X and a maximum of 10% of missing data per individual and variant.

### Population structure analyses considering the global dataset

A PCA was performed to explore the level of population structure over the five western Pacific BABs (Figure 2, A). This analysis showed a very clear geographical separation with two distinct clusters, one corresponding to the Manus and Woodlark basins and the other to the North-Fiji, Futuna and Lau basins. The first component (PC1) explained 26.03% of the total variance; the second one carried only 0.03% of the total variance (Figure 2 and SI Figure 8). This pattern was

consistent with the AMOVA results (Table 1), which also showed a strong and significant genetic differentiation between Manus/Woodlark and North-Fiji/Futuna/Lau, but no differentiation among basins and localities within these two groups (between Manus/Woodlark and North-Fiji/Futuna/Lau,  $F_{ST} = 0.387$ , *p*-value < 0.001, inter-basins within Manus/Woodlark and North-Fiji/Futuna/Lau,  $F_{CT} = -0.050$ , NS). In addition, the between-basins pairwise  $F_{ST}$  (Table 2) were only significant between Manus/Woodlark and North-Fiji/Futuna/Lau pairs. No significant pairwise  $F_{ST}$  values were observed between localities within either of the two groups Manus/Woodlark and North-Fiji/Futuna/Lau (SI Table 3).

This finding is also consistent with the ADMIXTURE (Figure 2 B & SI Figures 9–10) clustering results and strongly supports the same two distinct clusters (K = 2) with very few individuals showing very low percentages of mixed ancestry (from 0.1% to 3%). The identity-by-state distribution (SI Figure 11) did not show evidence of any internal structure due to kinship.

TreeMix analyses showed an optimal number of two migration events, whereas additional events did not improve the likelihood (SI Figure 12 A). Displaying the first migration event showed a very low migration weight from Manus/Woodlark towards North-Fiji (SI Figure 12 B), and adding the second migration event indicated a very slight differentiation between Woodlark and Manus (SI Figure 12 C). The  $F_3$  statistics showed a significant admixture signal, with source populations from each genetic cluster only when North-Fiji was chosen as the focal population (SI Figure 13).

The estimated genetic diversity of the populations considering all DNA positions of the 10 570 ddRAD-tags was bimodal, with slightly higher nucleotide diversity in Manus/Woodlark ( $\pi = 0.00572$ ) compared with North-Fiji/Futuna/Lau ( $\pi = 0.00535$ ), regardless of the populations being considered by geographic basin or by genetic cluster (SI Figure 14). The raw nucleotide divergence ( $D_{xy}$ ) between the two genetic clusters was estimated to be 0.0136.

Hence, the analyses of the complete SNP dataset of *I. nautilei* indicate the co-occurrence of two quasi-panmictic metapopulations, one associated with the Manus/Woodlark basins and the other with the North-Fiji/Futuna/Lau basins, on either side of the Solomon Islands/New Hebrides arc. Thus, these two metapopulations are both sufficiently homogeneous and differentiated from each other to be analyzed using  $\partial a \partial i$  demo-genetic inference, which aims at summarizing the global genome-wide history of divergence/contact between them over a long period of time.

Inference of demographic history and gene flow

### 1.Model comparisons

The folded JAFS in Figure 3 (A) shows how allele frequencies are shared between the Manus/Woodlark and North-Fiji/Futuna/Lau metapopulations. The  $\partial a \partial i$  framework can fit population models on the observed dataset and compares them based on their AIC values (Figure 4).

Among the four simplest models (SI, AM, IM and SC), SC was significantly the best fitting model. Increasing complexity by adding the parameters G, 2m, 2N independently improved the AIC values regardless of the basic model used. However, capturing the effect of linked selection (2N) explained the data much better than models with population growth (G) and heterogeneous gene flow (2m) only (Figure 4).

Conversely, the combination of these parameters led to only a slight improvement in the AIC values. Nevertheless, models including the effect of linked selection (2N) were still better than the other models (Figure 4).

Hence, considering all models together, the best ones were those that took all parameters (2N, 2m, G) into consideration, followed by models with only 2N+G (Figure 4). Moreover, for the 2N+2m and 2N+2m+G models, the proportion 1-P of the genome that evolves under restricted migration in 2m models amounted to 0.52–0.56 for the best AIC simulation among all runs, meaning that the proportion of the genome that evolves under a reduced effective migration rate (barrier loci) may be quite substantial. With the increasing number of population parameters, the secondary contact hypothesis was no longer the best evolutionary scenario explaining our genetic dataset: the models IM+2N+2m+G, SC+2N+2m+G, AM+2N+2m+G and AM+2N+G, IM+2N+G, SC+2N+G showed very similar AICs ( $\Delta_{AIC} \leq 10$ , Figure 4).

### 2. Inferences of model parameter values

According to the best models based on AIC, the two metapopulations may have diverged due to early (AM), late (SC) or constant (IM) gene flow and it is difficult to distinguish among these three possibilities. However, these models have some interesting features in common. First, although the standard deviations (SDs) are rather large, the effective population sizes of the two derived populations estimated since the split (N<sub>1</sub> & N<sub>2</sub>) indicate a demographic expansion (b1 and b2 > 1), regardless of the model, including a temporal change in population size (G). Second, the

local effect of selection at linked sites seems to affect a very large proportion of loci (Q = 0.99) with a small value of *hrf* (Hill-Roberston factor = ~ 0.02) (Table 3). Third, the number of contemporary migrants (estimated by  $(N_1*b1*m_{12})/2$  and  $(N_2*b2*m_{21})/2$ ) shows asymmetrical, but weak flow between the two metapopulations, slightly higher from North-Fiji/Futuna/Lau to Manus/Woodlark (4.2–4.6) than in the opposite direction (2.9–3.2). Fourth, nearly half of loci show a restricted migration rate (0.52  $\leq$  1-P  $\leq$  0.56). Fifth, the total estimated divergence time expressed as the number of generations are very similar, regardless of the model (Tsm, Tam+Ts, Ts+Tsc) and estimated to be between 66 951 and 70 295 generations.

### Outlier loci and detection of fine-scale structure

Despite the absence of geographic structuring within each metapopulation depicted in the global analysis, several outlier loci were identified in each metapopulation at the thresholds of  $p \le 0.05$  and  $p \le 0.01$  (SI Table 4).

BayeScan identified much fewer candidate outliers than expected by chance only, and hence was not considered further. PCAdapt was largely out of its working range because it searches for loci that exceed the possible differentiation level captured by the very first principal component as opposed to the second-order axes. However, all axes, except axis 1 which separates the two geographic metapopulations, belong to this second category due to the lack of internal structure. Hence, all the second-order axes primarily captured noise, and they were unable to reveal additional structuring (SI Figure 15,16). Therefore, Baypass and Arlequin were the only two methods considered further. To increase the probability of considering true positives only, we only kept loci identified in both approaches (predicted by the intersection depicted in the Venn diagram in Figure 5 at the thresholds  $p \le 0.05$  and  $p \le 0.01$  in SI Figure 17). Only 458 and 223 outlier loci were shared between the methods Baypass and Arlequin in Manus/Woodlark and North-Fiji/Futuna/Lau at the threshold of  $p \le 0.05$  respectively.

PCA based on these different sets of outlier loci helped to visualize their contribution to the internal heterogeneity of each metapopulations (Figure 5 and SI Figure 18). Interestingly, although outliers were defined within each metapopulation, the inter-metapopulation differentiation was retrieved in all cases. Nevertheless, a clear signal of differentiation was highlighted within both regions. Individuals from the Manus and Woodlark basins showed clear genetic differentiation with no overlap on PC2 (Figure 6 A). Individuals from North-Fiji were slightly pulled towards

Manus/Woodlark individuals based on PC1 (Figure 6 B), but they were also shifted on PC2 when outliers were considered at the threshold of  $p \le 0.05$ .

The ADMIXTURE analyses based on the outlier SNPs datasets with the threshold of  $p \le 0.05$  displayed optimal *K* values at *K*=2. With North-Fiji/Futuna/Lau outliers, North-Fiji displayed an admixture proportion from Manus/Woodlark ranging between 5 and 15% (Figure 6 C). For Manus/Woodlark outliers, we also found an admixture proportion from North-Fiji/Futuna/Lau in Woodlark (Figure 6 D). However, very similar values of cross-validation errors were obtained with *K*=3 (SI Figure 19,20,21), North-Fiji and Woodlark each being individualized as the third cluster in their respective runs.

The  $F_3$  statistics calculated based on outliers only did not provide any additional information (SI Figure 22).

When blasted onto the *A. boucheti* transcriptome, 30% of outlier loci identified at the threshold of  $p \le 0.05$  matched with the coding sequences of transcribed regions. This number was greater than expected by chance from randomly picked ddRAD loci along the *I. nautilei* genome. Half of these 30% of outlier loci (129 transcripts) had annotations on the protein database. Among these annotations, many involved genes that encode for DNA/RNA replication and repair enzymes, transmembrane carriers and synapse/microtubule biosynthesis, but also genes involved in the exocytosis/endocytosis regulation, and more especially the GTPase regulation pathway (SI Table 7). In addition, two genes involved in spermatogenesis were also detected.

### Discussion

Previous work by Thaler et al. (2011) using microsatellites and mitochondrial *cox1* sequences demonstrated that the southwestern Pacific deep-sea hydrothermal vent gastropod *Ifremeria*. *nautilei* is genetically structured into two distinct populations from Manus and North-Fh-Fiji/Lau BABs. Our study extends these previous results to a finer scale, owing to our larger, nested sampling design that includes the newly discovered La Scala vent field in the Woodlark basin

(Boulart et al., in press), the Futuna volcanic arc (Konn et al., 2016) and the newly discovered northernmost Mangatolo site at the entrance of the Lau basin. Using a 10 570 SNP genome-wide dataset—unprecedented for a hydrothermal species—, we confirm that *I. nautilei* is structured into two loosely connected metapopulations corresponding to two BAB ensembles. These ensembles display an almost complete internal genetic homogeneity; however, our analyses of outlier loci nevertheless revealed fine-scale differentiation among basins within each metapopulation. We discuss below the possible implications of these results in terms of larval dispersal and demographic connectivity and ultimately their consequences on the resilience of hydrothermal communities.

#### Long-term gene flow and history of differentiation

One metapopulation comprises the Manus and Woodlark basins (i.e. the Manus/Woodlark BAB) west of the Salomon/New Hebrides arc, whereas the other extends east of it with the North-Fiji basin, Futuna volcanic arc and Lau basin (i.e. the North-Fiji/Futuna/Lau BAB). The genetic divergence between the two metapopulations is relatively strong (average  $F_{ST} = 0.387$ ,  $p \le 0.001$ ,  $D_{xy} = 0.0136$ ), but each of these two ensembles appears to be pannictic ( $F_{CS} = -0.05$ , NS, SI Table 3). The demo-genetic inferences gleaned from  $\partial a \partial i$  suggest that the two metapopulations diverged with only a brief period of isolation (Ts was found to vary between 0.001 and 0.443 in the AM and SC models), although the existence of constant gene flow (IM) could not be formally excluded. The incorporation of several other demographic parameters (2N, 2m, G) produced a clear improvement in model fit. Considering each parameter independently, the effect of linked selection (2N) had a much greater influence on AIC than the other two parameters (2m and G), suggesting that a non-negligible proportion of loci may be influenced by linked selection. For the best models (2N+2m+G), this proportion approaches Q = 0.99 (which seems to be unrealistic), whereas only half of the markers appear to be under the influence of heterogeneous migration ( $0.52 \le 1-P \le 0.56$ ). Nevertheless, these models are very close to the 2N+G models ( $\Delta_{AIC} \le 10$ ), which estimate a proportion of loci under linked selection ( $0.56 \le Q \le 0.58$ ) and do not take into account the effect of heterogeneous migration. Disentangling these two effects is thus difficult and suggests that there are many genomic regions, possibly with lower recombination rates, where background selection and possibly selective sweeps have accelerated the rate of lineage sorting during divergence (Rougeux et al., 2017). This strong bimodality between two classes of loci affected or not by linked selection is also captured by the distribution of  $F_{ST}$ , which shows a clear

trough and then a peak around 0.15-0.2 (SI Figure 23). However, this bimodality reduces the ability to distinguish between the isolation-with-migration, the secondary contact, or the ancient migration scenarios in the more complex models (IM+2N+2m+G, AM+2N+2m+G and SC+2N+2m+G).

Considering an average DNA mutation rate of 10<sup>-8</sup>/site/generation, we estimated the time for the onset of divergence between the two metapopulations to be 60 000-70 000 generations (but admittedly this could as well be 10 times greater if the mutation rate is 10 times smaller). The generation time of *I. nautilei* is still unknown. Nevertheless, most hydrothermal species display an r-strategy suggesting short generation times (1-2 years) as an adaptation to the unstable and ephemeral nature of their habitat (Tyler & Young, 1999). Hence, we can suppose that the two populations started to diverge between 60 and 140 thousand ago (kya) for a mutation rate per site and per generation of 10<sup>-8</sup> and 10 times more with a mutation rate of 10<sup>-9</sup>. However, these estimates correspond to discrete non-overlapping generations and the reproduction of older cohorts may increase the equivalent generation time and, as a result, the divergence estimates. These values may be tentatively compared with estimates from the *cox1* sequences in Boulart et al. (in press) (net divergence 0.615% estimated on all sites). This latter value would amount to  $\sim$ 0.550 million years ago (*mya*) considering the widely used divergence rate of 1.4%/million years (mvr) for mitochondrial DNA (Knowlton & Weigt, 1998), but can reach 1.2 mya, depending on the average mitochondrial substitution rate considered for vent species (0.2-0.3%/myr (Chevaldonné et al., 2002; Breusing et al., 2020; Castel et al. in prep.). Although these estimates are notoriously highly variable and error-prone (see for instance Breusing et al., 2020), divergence time could range between 0.5 and 1 mya. This estimate is in rough agreement with Martinez & Taylor, (1996) who showed that the center of the Manus BAB started to spread quite recently (~ 0.78 mya), suggesting that hydrothermal vents within the spreading center may be younger than this estimate. Thus, it cannot be excluded that the divergence history of *I. nautilei* is relatively recent and not linked to the formation of BABs, but instead to regional modifications of surface and deep-sea currents during previous glacial maxima in relation to the extension of the Antarctic ice sheet which culminated around 0.126 mva (Barrows et al., 2011; Joy et al., 2014).

In addition to these divergence time estimates, the models allowed us to quantify the existence of an ongoing bidirectional and asymmetrical gene flow, with migration from North-Fiji/Futuna/Lau to Manus/Woodlark being higher than in the opposite direction. Despite this slight asymmetry, a genetic influence of the Manus/Woodlark metapopulation was detected in North-Fiji, which shows foreign alleles coming from the former rather than from the Lau/Futuna populations (also observed at mtDNA cox1 gene in Thaler et al., 2011 and Boulart et al., (in press), but not the other way around (*i.e.* influence of North-Fiji/Futuna/Lau on Woodlark, but see below). This result is consistent with the geography of the region, because North-Fiji and Woodlark are the closest BABs between the two metapopulations. Connectivity through larval dispersal between these two BAB ensembles has been tested by Mitarai et al. (2016) who simulated larval dispersal through entrainment of particles by oceanic currents prevailing at depths of 1000 m in the western Pacific. That study inferred a weak stepping-stone connection through a long planktonic larval duration (PLD of 170 days), provided that active hydrothermal sites in the Solomon and New Hebrides/Vanuatu arcs act as a relay. Such fields are known to exist, mostly associated with seamounts such as Nifonea, Tinakula or Stanton along the New Hebrides/Vanuatu arc (McConachy et al., 2005; Schmidt et al., 2017). The larval dispersal model developed by Mitarai et al. (2016) suggests a scenario where dispersal is mainly oriented from east to west: a situation also depicted in this region by Yearsley & Sigwart (2011) for a non-hydrothermal species at several depths (800 and 1400 m) and with various PLD lengths (27-151 days). However, when looking at surface countercurrents between Manus/Woodlark and North-Fiji/Futuna/Lau, Ganachaud et al. (2014) found surface currents oriented mainly from west to east through Solomon Islands and Vanuatu waters.

Cases of asymmetrical bidirectional gene flow between two metapopulations have also been found in two other hydrothermal gastropod species occurring sympatrically with *I. nautilei*, *L. schrolli* (Plouviez et al., 2019) and *A. boucheti* (Breusing et al., 2021). But, in contrast to *I. nautilei*, the predominant gene flow is oriented eastward from Manus to Lau, ( $m_{M->L} = 0.625$ ,  $m_{L->M} = 0.1725$ for *L. schrolli* and  $m_{M->L} = 12$ ,  $m_{L->M} = 2.6$  for *A. boucheti*).

Similarly to *I. nautilei*, *L. schrolli* is considered to possess lecithotrophic larvae (Berg, 1985; Craddock, Lutz, & Vrijenhoek, 1997; Tyler et al., 2008)). As for *A. boucheti*, its larval stage remains unknown, although both its morphology (Warèn & Bouchet, 1993) and the eDNA detection of *Alviniconcha* larvae close to the surface suggest planktonotrophy (Sommer et al., 2017). Provided that our estimates reflect ongoing migration, we hypothesize that *I. nautilei* larvae may be influenced by deep as well as surface currents, which could explain bidirectional gene

flow, one direction being slightly stronger than the other. This asymmetry suggests vertical migration of larvae. However, further investigations including oceanographic modeling and laboratory experiments are needed to address this hypothesis. For example, larvae of the hydrothermal gastropod *Shinkailepas myojinensis* (Yahagi et al., 2017) are able to migrate through the water column, and there is evidence of hydrothermal species' larvae in near-surface waters (Arellano et al., 2014, Sommer et al., 2017). Nevertheless, although many unknowns remain, our results indicate that *I. nautilei* has a complex dispersal strategy and pattern.

### Fine-scale population structure and connectivity

The high homogeneity of the two clearly distinct *I. nautilei* metapopulations necessarily entails that the intra-metapopulation migration (*i.e.* inter-localities within each BAB and inter-BABs within each metapopulation) is strong or extremely recent. Moreover, no kinship-related structure was detected in the SNP dataset, indicating that there is either no self-recruitment even though females brood their larvae to the trochophoran stage, or that population sizes are so large that the probability of detecting potential kin is too small (Table 4). Consequently, genetic connectivity within each metapopulation appears to be high, with evenly distributed polymorphisms among sampled sites despite the patchy distribution of hydrothermal vents and the inter-site distances, which may vary from hundreds of meters to more than a thousand kilometers within each metapopulation. This genetic connectivity therefore suggests that *I. nautilei* larvae are able to disperse within the range of each metapopulation after spawning.

The question is now whether this genetic homogeneity of each metapopulation arises from demographic connectivity (*i.e.* recruitment at one site being strongly influenced by the exportation of propagules from other sites) or is due to sporadic/rare larval exchanges able to counterbalance very attenuated genetic drift due to large local population sizes. The mechanism behind the observed genetic homogeneity has strong implications in terms of conservation biology, because demographic connectivity can play a crucial role in the resilience of populations faced with local extinction potentially exacerbated by deep-sea mining. The global analysis relying on a panel of primarily neutral markers indicates no differentiation at the metapopulation scale, but—as advocated by Gagnaire et al., (2015)— a few loci markers potentially undergoing direct, or indirect selective pressures may locally harbor distinct allele frequency in the recipient population. This pattern can be explained by local selection for foreign alleles that are better adapted or less loaded by deleterious mutations than resident ones, or by resolving intrinsic asymmetrical

incompatibilities between divergent genomes (Simon et al., 2021) creating local soft sweeps through linked selection. These processes result in enhanced local introgression of certain marker loci, a common pattern observed in blue mussels (Fraïsse et al., 2016) or European sea bass (Robinet et al., 2020), for example. These markers will appear as  $F_{\rm ST}$  outliers that may indicate recent dispersal events.

Our outlier analyses indeed suggest introgression of some loci. In Figure 6 B and SI Figure 18 B, individuals from the North-Fiji basin seem to be closer to Manus/Woodlark than Lau/Futuna on PC1, which may correspond to introgression in some of the outlier loci. An introgression pattern was confirmed by the  $F_3$  tests performed with the North-Fiji basin as the focal populations (significant negative value of the  $F_3$  statistic, SI Figure 13) and the ADMIXTURE analyses (Figure 6 C). These results indicate that some alleles found at high frequency in North-Fiji individuals are the consequence of long-range migration from Manus/Woodlark. Interestingly, with Manus/Woodlark outliers, although not visually detectable on the PCA (Figure 6 A and 18 A), Woodlark individuals exhibit some level of admixture from the North-Fiji/Futuna/Lau metapopulation (Figure 6 D). This low-level admixture corroborates our inference of ongoing bidirectional gene flow. However, it is not yet clear as to why its impact appears stronger in populations of the North-Fiji basin, against the predominant direction according to our  $\partial a \partial i$ inferences. Although we are unable to determine the precise mechanisms behind these frequency changes, these alleles have not diffused from North-Fiji to Lau/Futuna, indicating a subtle-but real-limitation in connectivity between the former and the latter. The same reasoning applies for the traces of admixture detected in Woodlark that appear to have not diffused to Manus.

Another kind of differentiation depicted by outlier loci seems to be explained by intrametapopulation divergence. The question arises as to the origin of these slight divergences on a PC axis orthogonal to the main inter-metapopulation divergence, which does not necessarily proceed from gene flow between differentiated populations as described above. Allele frequency differences for outlier loci between Manus and Woodlark are detectable on PC2 (Figure 6 A, SI Figure 18 A, ADMIXTURE K = 3, SI Figure 20). The same question applies to the eastern North-Fiji/Futuna/Lau metapopulation, with differences between North-Fiji and Futuna/Lau (Figure 6 B, ADMIXTURE K = 3, SI Figure 21). This pattern can be due to any combination of drift and/or selection. Local selection may result from major differences in depth or vent fluid composition. The fact that the fraction of outliers mapping on transcribed regions is greater than by chance and targets a few metabolic/regulatory pathways suggests their possible involvement in local adaptation to depth or different fluid chemistry, but this remains to be investigated (Jennings et al., 2013; Liu et al., 2021). In the absence of high demographic connectivity required to ensure the interdependency of local populations, this local differentiation can remain detectable for several generations before being shuffled among all metapopulation demes.

These subtle limitations in connectivity between basins can be associated with abyssal plains, which may limit gene flow in a disconnected ridge system such as that found at the regional scale of these BABs. Physical barriers in other parts of the world, such as transform faults and microplates, have already been shown to greatly impede the effective migration of deep-sea vent species at a much more restricted spatial scale (Johnson et al., 2006; Plouviez et al., 2009; Plouviez, et al., 2013). However, regarding the populations of the Futuna volcanic arc and Lau basin, our in-depth scrutiny of outliers did not reveal any sign of genetic differentiation. Hence, the hypothesis of demographic correlation between these two regions cannot be rejected, although we cannot infer with certitude the directionality of the exchanges.

### Conclusions

Overall, our analyses revealed a clear genetic differentiation of *Ifremeria nautilei* populations between the Manus/Woodlark and the North-Fiji/Futuna/Lau BABs, with very high gene flow within each of these two metapopulations as well as higher genetic diversity in Manus/Woodlark. Despite an in-depth scrutiny of genome-wide genetic variation, no geographic substructure was detected between or within localities sampled within each individual ridge system. This genetic connectivity probably indicates high local (re)colonization capacity for this hydrothermal vent species due to the ephemeral nature of active sites in this region, at least at the scale of a given back-arc basin.

However, our outlier analyses revealed that this genetic connectivity does not necessarily equate with demographic connectivity at the larger inter-basin intra-metapopulation scale. The specific investigation of outlier loci illustrates how a few loci in a large genome-wide dataset can carry useful information about actual barriers to dispersal in high gene flow species. Deep-sea mining holds the potential to exacerbate dispersal barriers and limit population resilience, because if a large proportion of the vent habitat is destroyed locally, population rescue from other basins will be restricted.

Furthermore, our demographic simulations indicated a long period of divergence during the Quaternary period (several tens of thousands of generations) associated with restricted long-range gene flow over a large fraction of the genome. Although the effects of linked selection and reduced migration (barrier loci) are not clearly distinguishable, our results suggest that the effect of the latter is less pronounced. This interpretation agrees with the fact that the global divergence among the two metapopulations is still quite low (net nuclear nucleotide divergence, 0.81%). This divergence perhaps reflects the very beginning of an ongoing speciation process, where a handful of barrier loci may already exist and at the same time overall genetic differentiation is not hampered by weak contemporary and asymmetrical gene flow between metapopulations.

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### **Data Accessibility**

### Genetic data:

Raw sequence reads (Individual fastq files) are available at the European Nucleotide Archive (study accession number PRJEB47533) https://www.ebi.ac.uk/ena/browser/view/PRJEB47533).

Individual genotype data (VCF) are available on DataDryad: https://doi.org/10.5061/dryad.ffbg79cwq

Scripts used in this study (R,  $\partial a \partial i$ ) are available on a public Github repository: (https://github.com/Atranluy/Scripts-Ifremeria).

### Sample metadata:

Metadata are also available on DataDryad: https://doi.org/10.5061/dryad.ffbg79cwq

### **Benefit-Sharing Statement:**

In order to obtained the requested authorizations to work in national waters and in agreement with the Nagoya protocol, we contacted the authorities of the different countries (Papua-New Guinea, Fiji, and Tonga) and territories (Wallis and Futuna) for benefit-sharing where sampling was performed. The data generated will be accessible on public databases. The results obtained will also be communicated to these authorities which may have to make decision regarding conservation of deep-sea hydrothermal vent communities in their EEZs. Observers for the different countries who took part of the on-board activities will be informed of our findings.

### Author's contribution:

D. J. and S. H. designed the CHUBACARC and CERBERUS projects, F. B. supervised the genetic work. A. T. L. Y., S. R., C. D. T., J. C., P. W. and A. P. performed laboratory work. A. T. L. Y. performed bioinformatics statistical analyses with the contribution of F. B., D. J., P. A. G., N. B. and T. B. A. T. L. Y., F. B. wrote the manuscript with feedback of T. B., D. J., P. A. G., N. B., S. A. H. and C. D. T. All authors approved the manuscript.

# Tables

Table 1: Analysis of molecular variance (AMOVA) on the final dataset with 10 000 permutations (\*\*\*: p < 0.001, \*\* : p < 0.01 \*: p < 0.05)

Manus/Woodlark vs. North-	0.38773***	$F_{\rm ST}$
Fiji/Futuna/Lau		
Basins in M/W and NF/F/L	-0.05	$F_{\rm CT}$
Localities within Basins	-0.00011	$F_{\rm SC}$
Individuals within Localities	-0.05084	$F_{ m IS}$

M/W: Manus/Woodlark, NF/F/L: North-Fiji/Futuna/Lau

Table 2: Pairwise (between basins)  $F_{ST}$  matrix on the final dataset with 10 000 permutations after Bonferroni correction (\*\*\*: p < 0.001, \*\*: p < 0.05).

		Lau	Futuna	North-Fiji	Manus	Woodlark
	Lau	0.00000				
	Futuna	-0.00040	0.00000			
1	North-Fiji	0.00029*	-0.00004	0.00000		
	Manus	0.38350***	0.38275***	0.37651***	0.00000	
	Woodlark	0.39986***	0.39656***	0.38647***	-0.00016	0.00000

Table 3: Parameters estimated from  $\partial a \partial i$  for the three best models (IM2N2mG, SC2N2mG and AM2N2mG\*) with their standard deviations (SD) estimated using a Fisher information matrix. (\*isolation with migration (IM), secondary contact (SC), ancient

migration (AM) and with parameters describing effective population size (2N), migration rate (2m) and population growth (between basins))

	Parameter	IM+2N+2m+G	SD	SC+2N+2m+G	SD	AM+2N+2m+G	SD
	N <sub>1</sub> (NF/F/L)	0.435	0.187	0.913	0.573	0.390	0.127
	N <sub>2</sub> (M/W)	0.411	0.157	0.840	0.573	0.356	0.119
	b1	30.410	13.176	16.947	8.127	34.367	12.399
	b2	25.097	8.683	13.714	8.451	29.288	10.523
	hrf	0.023	0.006	0.021	0.006	0.022	0.006
	Ts			0.443	0.527	0.001	0.024
	Tsm/Tsc/Ta m	1.631	0.280	1.470	0.379	1.681	0.335
	m12	0.444	0.147	0.422	0.145	0.461	0.137
_	m21	0.825	0.192	0.810	0.261	0.825	0.198
	me12	0.038	0.020	0.038	0.024	0.039	0.018
	me21	0.283	0.063	0.270	0.088	0.300	0.070
	Р	0.483	0.103	0.439	0.093	0.471	0.095
	Q	0.990	0.136	0.990	0.185	0.990	0.136
	Theta	271.772	31.55	243.283	64.91	273.222	32.988

N represents the population size of each population; b, the population growth factor; hrf, the Hill-Robertson factor; Ts, the time of strict divergence; Tm/Tsc/Tam, the time of divergence with migration; m12, represents the unrestricted migration rate from the population 2 towards population 1; me12, the restricted migration rate (e.g. barrier loci) from population 2 towards population 1; Q, the proportion of loci that are under the effect of linked selection (i.e. Hill-Robertson effect); P, the proportion of loci that have unconstrained migration; M/W, Manus/Woodlark; NF/F/L, North-Fiji/Futuna/Lau

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Table 4: Estimates of the effective number of migrants( $N_m$ ) exchanged per generation between metapopulations, total time of divergence since the population split and effective population size ( $N_e$ ) for three demographic models (isolation with migration (IM), ancient migration (AM), and secondary contact (SC).

	N <sub>m</sub> 2->1	N <sub>m</sub> 1->2	Т	$N_1$	$N_2$
	(M/W->NF/F/L)	(NF/F/L-> M/W)	(in generations)	(NF/F/L)	(M/W)
IM+2N+2m+G	2.935	4.255	66 951	271 506	211 708
SC+2N+2m+G	3.265	4.665	70 295	284 279	211 653
AM+2N+2m+	3.09	4.305	69 380	276 561	215 142

Metapopulation M/W, Manus/Woodlark; metapopulation NF/F/L, North-Fiji/Futuna/Lau.

# **Figures**

Figure 1: Colors: Back-arc basins. Sampling map of Ifremeria nautilei in the Southwestern Pacific Ocean. The number of localities is given in parentheses and the total number of sampled sites in brackets.

Figure 2: (A) Principal component analysis plot of 362 Ifremeria nautilei individuals from five hydrothermal basins scored at 10 570 SNPs, open circles represent the multivariate normal distribution of each group (basins) at 95%. (B) ADMIXTURE plot for each individual with their ancestry proportions obtained on the final dataset for the best K (K=2). Individuals are grouped according to their basin of origin.

Figure 3: (A) Joint allele frequency spectrum (JAFS) between the Manus/Woodlark (M/W) and North Fiji/Futuna/Lau (NF/F/L) basin systems. (B) Simulated JAFS under the IM2N2mG model (see Figure 4), the log scale indicates the density of SNPs in each frequency class. (C) Residuals of the fit of the simulated model on the data. (D) Representation of the fitted model. (N represents population size; b, population growth factor; hrf, the Hill-Robertson factor, which simulates linked selection; m, unrestricted migration rate; me, restricted migration rate, which simulates barrier loci)

Figure 4: Heat-map of the best Akaike information criterion (AIC) value for each parameter combination (population expansion or contraction (G), effect of linked selection (2N) and heterogeneous migration (2m)) and demographic model (strict isolation (SI), isolation with migration (IM), ancient migration (AM), and secondary contact (SC)).

Figure 5: Venn diagram of shared outlier loci identified in Arlequin and Baypass with a p-value less than or equal to 0.05 within each metapopulation (Manus/Woodlark, M/W and North Fiji/Futuna/Lau, NF/F/L) independently.

Figure 6: Principal component analysis on all individuals with the outlier loci found in both Arlequin and Baypass at a 0.05 p-value threshold. (A) Outlier loci detected in Manus/Woodlark (M/W). (B) Outlier loci detected in North-Fiji/Futuna/Lau (NF/F/L). Plot of ancestry proportion inferred with Admixture at K = 2 on all individual at the 0.05 p-value threshold, for (C) North-Fiji/Futuna/Lau outliers and (D) for Manus/Woodlark outliers.









Simple-	2278.94	2276.73	2169.45	3167.78	
G-—	1853.05	1864.15	1834.24	2930.87	
2N+G -	1330.51	1331.61	1333.81	1917.85	AIC Value
ameter 52N+2m+G -	1326.66	1325.12	1326.78		3000 2500
2N+2m -	1376.42	1370.42	1347.3		2000 1500
2N-	1351.12	1349.11	1341.59	1963.73	
2m+G -	1372.76	1370.88	1373.96		
2m-	1674.98	1741.13	1648.9		
	AM	SI			



