RESEARCH ARTICLE

Considering reefscape configuration and composition in biophysical models advance seascape genetics

Simon Van Wynsberge1,2,*, Serge Andréfouët3, Nabila Gaertner-Mazouni1, Josina Tiavouane2, Daphné Grulois2, Jérôme Lefèvre3, Malin L. Pinsky4,5, Cécile Fauvelot2

1 UMR-241 EIO, Université de la Polynésie Française, Laboratoire d'Excellence CORAIL, Faa’a, Tahiti, French Polynesia, 2 UMR-9220 ENTROPIE (Institut de Recherche pour le Développement, Université de La Réunion, CNRS), Laboratoire d’excellence-CORAIL, centre IRD de Nouméa, Nouméa, New Caledonia, 3 UMR-065 LEGOS and UMR-235 MIO, Centre IRD de Noumèa, Noumèa, New Caledonia, 4 Department of Ecology, Evolution, and Natural Resources, Rutgers University, New Brunswick, New Jersey, United States of America, 5 Institute of Earth, Ocean, and Atmospheric Sciences, Rutgers University, New Brunswick, New Jersey, United States of America

* simon.vanwynsberge@gmail.com

Abstract

Previous seascape genetics studies have emphasized the role of ocean currents and geographic distances to explain the genetic structure of marine species, but the role of benthic habitat has been more rarely considered. Here, we compared the population genetic structure observed in West Pacific giant clam populations against model simulations that accounted habitat composition and configuration, geographical distance, and oceanic currents. Dispersal determined by geographical distance provided a modelled genetic structure in better agreement with the observations than dispersal by oceanic currents, possibly due to insufficient spatial resolution of available oceanographic and coastal circulation models. Considering both habitat composition and configuration significantly improved the match between simulated and observed genetic structures. This study emphasizes the importance of a reefscape genetics approach to population ecology, evolution and conservation in the sea.

Introduction

The dispersal of individuals between populations is the foundation of meta-population dynamics [1]. In a meta-population context, dispersal can affect population growth and vital rates, a process termed “demographic connectivity” (Table 1) [2]. Characterizing the patterns of demographic connectivity and identifying their drivers is necessary for designing effective management plans [3, 4]. In practice however, measuring dispersal for many species at scales relevant to management questions is a difficult task [5]. Demographic connectivity is therefore often estimated with models but rarely validated against empirical field data [4].

Recent advances in population genetics and the identification of hypervariable genetic markers provide new opportunities to infer demographic connectivity by estimating gene flow
among populations from allele frequencies [2]. Indeed, it is often assumed that processes driving dispersal and demographic connectivity also shape genetic patterns [12]. Genetic connectivity is the degree to which gene flow affects evolutionary processes within populations [2]. However, genetic connectivity integrates dispersal over several generations, as well as other evolutionary forces (i.e., genetic drift, selection and mutation).

Trying to relate genetic connectivity with environmental factors expected to influence dispersal has been the focus of several recent studies. This approach was termed “landscape genetics” in terrestrial ecosystems, by a fusion of “landscape ecology” with “population genetics” [13]. Landscape genetics explicitly aims to quantify the effects of landscape composition and landscape configuration on gene flow and spatial genetic variation [6]. In the landscape ecology literature, the “landscape” is conceptualized as a complex of habitat patches. The term “landscape composition” refers to the nature and relative proportion of habitat patches in the landscape, while “landscape configuration” refers to the spatial arrangement of habitat patches relative to each other [7].

The simplest form of landscape genetics relates genetic distances to geographical distances (i.e., Euclidean distance), a model termed “isolation by distance” (IBD). Isolation by distance is the process by which geographically restricted gene flow and random genetic drift generate genetic differentiation among populations on different habitat patches [8]. In terms of metapopulation dynamics, IBD suggests that the exchange of individuals between habitat patches decreases as the geographical distance between them increases. Besides geographical distance, other environmental features related to landscape composition and configuration have also been used to explain genetic patterns, including habitat fragmentation, topography, snow depth, and presence/absence of rivers [14]. In these cases, unfavourable environmental features between habitat patches become barriers to gene flow, following an Isolation by Barrier (IBB) model.

### Table 1. Definitions for the main technical terms used in this study, with references.

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
<th>References</th>
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<tbody>
<tr>
<td>Demographic connectivity</td>
<td>Process by which dispersal of propagules affects population growth and vital rates</td>
<td>[2]</td>
</tr>
<tr>
<td>Genetic connectivity</td>
<td>Degree to which gene flow affects evolutionary processes within populations</td>
<td>[2]</td>
</tr>
<tr>
<td>Landscape genetics</td>
<td>Field of study that explicitly quantifies the effects of landscape composition and landscape configuration on gene flow and spatial genetic variation</td>
<td>[6]</td>
</tr>
<tr>
<td>Landscape composition</td>
<td>Nature and relative proportion of habitat patches that shape the landscape</td>
<td>[7]</td>
</tr>
<tr>
<td>Landscape configuration</td>
<td>Habitat fragmentation and spatial arrangement of habitat patches relative to each other</td>
<td>[7]</td>
</tr>
<tr>
<td>Isolation by distance (IBD)</td>
<td>Process by which geographically restricted gene flow and random genetic drift generate genetic differentiation of populations on different habitat patches, resulting in a pattern of increasing genetic differentiation as a function of distance</td>
<td>[8]</td>
</tr>
<tr>
<td>Multi-disciplinary seascape genetics</td>
<td>Modelling approach that integrates biophysical information to adequately explain the observed population genetic structure in the marine realm</td>
<td>[9]</td>
</tr>
<tr>
<td>Isolation By Oceanographic Distance (IBOD)</td>
<td>Process by which genetic differentiation among populations is induced by the direction and intensity of water currents</td>
<td>[10]</td>
</tr>
<tr>
<td>Habitat patchiness</td>
<td>Level of fragmentation of habitat patches, from highly fragmented to continuous</td>
<td>[11]</td>
</tr>
<tr>
<td>Seascapes (as currently perceived in seascape genetics)</td>
<td>Complex mosaic of pelagic habitat patches. Pelagic habitat refers to environmental features of the water column.</td>
<td>-</td>
</tr>
<tr>
<td>Reefscape</td>
<td>Complex mosaic of benthic habitat patches in coral reef ecosystems.</td>
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<tr>
<td>Reefscape genetics</td>
<td>Field of study that explicitly quantifies the effects of reeascape composition and reascape configuration on gene flow and spatial genetic variation. While seascape genetics mostly focused on the pelagic habitat, reeascape genetics emphasis the role of benthic habitats on genetic structure.</td>
<td>This study</td>
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https://doi.org/10.1371/journal.pone.0178239.t001
The translation of landscape genetics to the marine realm was termed “seascape genetics” [9, 15], but has become its own fairly distinct field due to inherent differences of the marine environment and marine organisms. In the marine realm, dispersal of most of organisms occurs during the pelagic stage of their life cycle (e.g., as larvae), so that population genetic structure is likely related to the transport pathways of propagules [16]. In this case, the dispersal kernel is influenced both by biological and physical processes affecting the pelagic larvae [4]. Approaches that integrate biophysical information on ocean current and larval dispersal have therefore become popular [9]. These “multi-disciplinary seascape genetics” approaches [9] introduced the concept of “Isolation By Oceanographic Distance” (IBOD), which relates empirical genetic structure with a metric of oceanographic distance as the seascape feature [10]. Information integrated into the distance metric often includes the Pelagic Larval Duration (PLD) and the direction and velocity of oceanographic currents [17, 18], though some studies have also investigated the influence of other environmental features of the water column that may act as barriers to the dispersal of propagules, including upwelling, gradients in water temperature, and salinity [16].

More recently, however, several modelling and demographic studies have suggested that seascape features beyond oceanographic distance may influence demographic connectivity. This includes, for example, the spatial configuration of favourable habitat for mature adults of benthic species (i.e., benthic habitat). In coral reef ecosystems, the spatial configuration of benthic habitat is referred to as the “reefscape configuration.” Indeed, habitat patchiness may drive the relative influence of local and non-local offspring, with higher self-recruitment expected in fragmented habitats compared to continuous habitat patches [11]. At time scales of several generations, habitat patchiness can also influence population genetic structure because distant habitat patches (which may not be connected directly by a single dispersal event) can nevertheless be indirectly connected by gene flow through intermediate habitat patches that act as stepping-stones. Stepping stone models therefore predict higher genetic structure in fragmented habitats than in continuous habitats [17]. In addition to habitat patchiness and reefscape configuration, reefscape composition is also likely to play a role in demographic connectivity. Habitat patches have to be of sufficient quality and suitability for a species to settle, survive, feed, grow and reproduce [19].

Despite these recent conceptual advances, few marine studies have tried to relate genetic structure to the spatial configuration and composition of adult habitats [20]. Improving our understanding of these links is a question of broad interest for ecology because spatial features related to reefscape may significantly influence population connectivity and functioning. In addition, a better understanding of reefscape influences on connectivity could help provide practical information for management and conservation where habitat mapping are available but larvae transport pathways are not.

The hypothesis to test in this study is that reefscape configuration and composition shapes genetic patterns. We propose explicitly quantifying the influence of reefscape configuration and composition on genetic structure with a multi-disciplinary modelling approach (Fig 1) that includes (i) establishing maps of suitable habitat for the focal species; (ii) estimating the relative probability of propagule dispersal between all habitat patches based on simple Euclidian distance or complex biophysical modelling; (iii) simulating gene flow over time to account for stepping stone processes as well as other evolutionary forces (e.g., genetic drift and mutation) and to translate the relative probability of propagule dispersal into genetic distances between all pairs of habitat patches; (iv) quantitatively comparing the simulated genetic structure with the observed genetic structure; (v) and performing sensitivity analyses to evaluate the extent to which the fit between simulated and observed genetic structure is affected when the original patterns of reefscape composition and configuration are modified or degraded.
Developing such an approach that integrates habitat suitability maps, biophysical modelling, gene flow modelling and empirical genetic data is, however, complicated by the range of spatial and temporal scales inherent to these processes [21]. To date, this has only been partially achieved in the seascape literature. D’Aloia et al. [22] investigated a method to infer the dispersal kernel of the Caribbean fish *Elacatinus lori* on the Mesoamerican Barrier Reef from genetic parental data while considering the spatial distribution of habitat patches. For the same species and in the same area, D’Aloia et al. [23] tested the influence of geographical distance (IBD) and oceanic breaks (IBB) on genetic structure and found that a 20 km break in the reef...
could induce genetic structure. Alberto et al. [24] modelled the spatial genetic structure of the giant kelp *Macrocystis pyrifera* obtained from microsatellite empirical data, as a function of geographic distance and habitat continuity (i.e. similar to habitat fragmentation). They found a significant influence of habitat continuity on genetic distance. Nanninga et al. [25] highlighted the effect of habitat quality on population genetic patterns of an anemonefish in the Red Sea, but their study considered habitat as the type of water masses surrounding the reefs and do not consider the benthic habitat of the targeted clown fish. Other studies integrated biophysical modelling in their analyses to test for IBOD: Davies et al. [18] tested both IBD and IBOD to explain the genetic structure and diversity of two corals (*Acropora hyacinthus* and *A. digitifera*) in Micronesia. They found a strong influence of geographic distance and oceanographic distance on genetic patterns and concluded that Micronesia may serve as a stepping stone pathway between the Coral Triangle in Asia (Philippines and Indonesia in particular) and the Central Pacific islands. Johansson et al. [26] investigated if oceanographic transport and other seascape features explained different scales of genetic structure of giant kelp, *Macrocystis pyrifera*. They found a significant effect of habitat continuity on genetic distance, but we are not aware of equivalent studies in coral reef ecosystems. Kool et al. [27, 28] used a matrix analysis in conjunction with a bio-oceanographic larval dispersal model to project the expected development of genetic structure in Caribbean and Indo-West Pacific coral reef ecosystems, but did not validate their model with empirical genetic data. Galindo et al. [29] used a coupled oceanographic-genetic model to predict population structure of Caribbean corals, and found a general concordance between the observed (from empirical data) and simulated genetic structure. They further highlighted that projecting connectivity forward in time provides a framework for studying long-term source-sink dynamics, making it possible to evaluate how dispersal can shape population genetic structure at regional scales [27, 28, 29]. While taking habitat maps as baseline for the oceanographic model, however, these authors did not explicitly test nor quantify the influence of reefscape composition and configuration on gene flow.

In this study, we followed the multi-disciplinary seascape modelling approach proposed above (see Fig 1) to test the hypothesis that reefscape configuration and composition shapes the population genetic structure of a common giant clam (*Tridacna maxima*) in New Caledonia and Vanuatu, South-West Pacific. The originality of our approach is about using a quantified, spatially explicit view of the benthic reef habitat of our targeted species coupled with the biophysical model. Specifically, we simulated the expected genetic diversifications among giant clam populations across the study area given the observed reefscape composition and configuration and under various scenarios of larval dispersal. These simulated genetic structures were compared to the genetic structure observed from empirical genetic data. Then, sensitivity analyses explicitly quantified the importance of habitat patchiness and habitat suitability on genetic structure. The giant clam *T. maxima* is our model species, but our results remain relevant for a wide range of species of similar life history traits (pre-competency period around 9 days and shallow reef habitat).

**Methods**

Permits for mantle biopsies of giant clams were from Direction de l’environnement de la Province Sud (n° 3117-2011/ARR/DENV, n° 2432-2012/ARR/DENV, n° 2660-2013/ARR/DENV), Direction du Développement économique et de l’environnement, Province Nord (n° 60912-25-28-2012/JJC, n° 60455-15-25/JJC), and Direction du développement économique de la Province des Iles Loyauté (n° 6161-37/PR). No permissions were required for Vanuatu.
Study area

The study area extended from 158°E to 171°E and from 24°S to 12°S. It covered Vanuatu’s oceanic islands and New Caledonia (Fig 2). Reefs of Vanuatu oceanic islands are mostly narrow fringing reefs with small, narrow lagoons. By contrast, a 1,600 km-long barrier reef surrounds both New Caledonia’s continental main island (Grande Terre) and its wide and deep 16,800 km² lagoon. Around Grande-Terre, several satellite reef systems of various sizes are found. The most prominent is further west: the “Chesterfield” archipelago includes a vast and open lagoon (12,200 km²) with several highly exposed intertidal reefs and islands [30]. The Loyalty Islands, Entrecasteaux atolls and Île des Pins reef systems are found respectively east, north and south of Grande Terre.

The main features and variability in regional circulation are reviewed and described in Cra-vatte et al. [32]. The westward South Equatorial Current (SEC) splits into several branches when crossing Vanuatu and New Caledonia, bifurcating either northwestward via the North Vanuatu Jet and the North Caledonian Jet, or southwestward via the South Caledonian Jet. Around the southwest area of New Caledonia, the surface flow is eastward via the Sub-Tropical Counter Current and the Alis Current of New Caledonia (ACNC), with high variability due to intermittent eddies. The Loyalty Islands are separated from the Grande Terre by the Vauban Current, usually oriented southeastward, but with high intra-seasonal variability. Indeed, transport pathways in the vicinity of Vanuatu and New Caledonia archipelagos are highly complicated as a result of offshore eddies migrating westward and local transient eddies and dipole circulations formed by interactions between bottom topography and ocean dynamics [32,33].

Reefscape composition and configuration

Habitat suitability maps were developed from field data on T. maxima densities and from 30 m Landsat satellite imagery from the Millennium Mapping Project [31]. Data on T. maxima densities came from Gilbert et al. [34], Wantiez et al. [35], and from Friedman et al. [36]. To avoid biases associated with combining data across multiple methods, we only considered studies that used belt transects. Overall, density estimates were available for 31 sites spread over the area. Each site was sampled with a number of stations spread over various shallow reef geomorphological units, and each station is a set of transects. Reef geomorphological units were defined at three levels (L1, L2, L3, Table 3 and Fig 3a and 3c). The geomorphological level L1 considered all reefs (shallow and variable depth reefs areas confounded) as habitat for T. maxima. By contrast, level L2 distinguished shallow reefs from variable depth reefs, the latter being characterized by hundred-fold lower densities of T. maxima. Since no quantitative information on giant clam density on variable depth reefs was available in the literature, we arbitrary fixed T. maxima density on variable depth reefs at the density reported in shallow reefs divided by one hundred. Level L3 dissociated “Outer barrier reef of continental islands” (found in New Caledonia) from “Intermediate reef of continental islands” (New Caledonia), “Fringing reef of continental islands” (New Caledonia), “Oceanic patch reef of continental islands” (New Caledonia), “Oceanic islands” (Vanuatu, Lifou, Mare and Tiga), and “Atolls/banks” (Entrecasteaux, Chesterfield). Densities of all L3 categories falling in the “Variable depth reefs” L2 category were fixed at the densities of the corresponding L3 category falling in the “Shallow reef” L2 category, divided by 100. For example, the density of giant clams in Shallow Fringing reef of continental island was estimated as 256 ± 272 ind.ha⁻¹ from field data, so we calculated the density of giant clams in Variable depth Fringing reef of continental island to be 2.56 ± 2.72 ind.ha⁻¹ (Table 3).
The continuous map of habitat was gridded at 0.25 × 0.25 degrees. Abundances of giant clam within each cell were calculated by multiplying estimated densities per habitat type by habitat area. The final grid was therefore a mosaic of habitat patches with different giant clam abundances (seascape composition) and different spatial relations to each other (seascape configuration) (Fig 3D).

**Dispersal kernel and connectivity matrices**

Two types of dispersal kernels were considered in this study. First, we tested the IBD model by considering the relative probability for larvae to travel from habitat patch \( i \) to habitat patch \( j \) \((P_{ij})\) as a decreasing function of geographic distance \( d \) (Eq 1):

\[
P_{ij} = a^d
\]

The continuous map of habitat was gridded at 0.25×0.25 degrees. Abundances of giant clam within each cell were calculated by multiplying estimated densities per habitat type by habitat area. The final grid was therefore a mosaic of habitat patches with different giant clam abundances (seascape composition) and different spatial relations to each other (seascape configuration) (Fig 3D).
A large panel of values was tested for \(a\) (from 0.40 to 0.99 by steps of 0.01), providing 60 scenarios of IBD. Geographical distance between habitat patches was defined as the shortest path to reach one cell from another without crossing land. Land maps for New Caledonia, Vanuatu, and Chesterfield were rasterized (0.1 degree resolution) using the rasterize function of the raster package in R 3.1.0, and the shortest path between each pair of cells was determined using the costDistance function of the gdistance package.

Second, we tested the IBOD model by considering the probability for larvae to travel from habitat patch \(i\) to habitat patch \(j\) as a function of oceanic currents. Larval dispersal between habitat patches was simulated with Lagrangian particle tracking models. We modelled the drift of larvae for 9 days, as that is the typical time for \(T.\ maxima\)’s Larval Precompetency Duration (LPD) [37, 38]. During the competency period (after 9 days), we allowed larvae to settle as soon as they crossed a patch of favourable habitat [39,40]. We considered a competency period ranging from 9 to 19 days on the basis of Jameson [37] and Neo et al. [40], with the number of surviving larvae exponentially decreasing with time. We used a survival curve similar to Wood et al. [41] (See S1 Fig).

The drift of particles through oceanic waters was simulated using the Roms Offline Floats (Roff, [42]) model based on a Regional Ocean Modeling System (ROMS, [43], see S1 File for details) with a 1/12’ (about 8 km) horizontal resolution that adequately reproduces mesoscale details of the mean flow [33]. Since no seasonality in recruitment is evident for \(T.\ maxima\) in New Caledonia, the rate of particle release was set to one per day from each cell from the top depth layer over the period 1993 to 2010. This was a period long enough to

<table>
<thead>
<tr>
<th>Location code</th>
<th>Location name</th>
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<tbody>
<tr>
<td>EFA</td>
<td>Efate</td>
<td>14</td>
</tr>
<tr>
<td>CHE</td>
<td>Chesterfield</td>
<td>19</td>
</tr>
<tr>
<td>HUO</td>
<td>Huon</td>
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<tr>
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<td>IP</td>
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</tr>
<tr>
<td>CS</td>
<td>Corne Sud</td>
<td>41</td>
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https://doi.org/10.1371/journal.pone.0178239.t002
Overall, $15 \times 10^5$ particles were released (6,500 per cell), which is expected to produce robust estimations of dispersal [44]. For each simulation, we identified the habitat patches where larvae were expected to settle considering oceanographic currents. This provided a dispersal matrix $D_{ij}$ containing the probability of transport for larvae between patches $i$ and $j$ through oceanic waters.

**Modelling gene flow**

To evaluate the expected genetic structure across the study area under a given hypothesis of larval dispersal, we modelled gene flow among populations with a matrix projection model following Kool et al. [27]. For a given locus and allele, we defined $Q_t$ as a 243 row matrix containing allele frequency among the 243 habitat patches (Eq 2). Model simulations started with an initial matrix $Q_0$ that contained frequency of alleles randomly (99 simulations) distributed among habitat patches.

$$Q_t = \begin{bmatrix} F_1 \\ \vdots \\ F_{243} \end{bmatrix}$$

(2)

The model then projected the allele frequency over time according to Eq (3):

$$Q_{t+1} = AQ_t + NQ_t$$

(3)
where $N$ is the number of individuals in each habitat patch, calculated from patch area and $T$. \textit{maxima} density per habitat. The density per habitat was randomly generated from a Gaussian distribution with mean and standard deviation estimated from our literature review (Table 3). $A$ is a square matrix (Eq 4) containing the number of migrants dispersing between pairs of

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Fig 3. Illustration of how habitat composition and configuration were taken into account in this study. (A), (B), and (C) are habitat maps for south BOU (see Fig 2) established according to reef typology levels L3, L2, and L1, respectively (see Table 3). (D) Estimated abundance of $T$. \textit{maxima} for each habitat patch. Abundance is the product of habitat surface and density per habitat (the figure displayed here was established on the basis of the L3 level, Table 3). Note that lower abundances were predicted in the south east part of New Caledonia and Vanuatu.
habitat patches from \( t \) to \( t+1 \) (i.e., the realized larval connectivity *sensu* Watson et al. [45]).

\[
A = \begin{bmatrix}
A_{1,1} & \ldots & A_{1,243} \\
\vdots & \ddots & \vdots \\
A_{243,1} & \ldots & A_{243,243}
\end{bmatrix}
\]

The interpretation of Eq 3 is that the expected state of the system \((Q)\) at \( t+1 \) is the result of migration progeny \((AQ_t)\) added to the original parent generation \((NQ_t)\). As \(Q\) refers to a matrix of allelic frequencies, a row normalisation among alleles was performed at each time step (indicated by top bar in Eq 3) so that the values always ranged between 0 and 1.

The number of migrants transferring from habitat patch \( i \) to habitat patch \( j \) \((A_{ij})\) during a=generation was considered time invariant and was calculated as the number of clams in habitat patch \( i \) \((N_i)\), multiplied by a fertility rate \( f \) (i.e., number of adult clams produced per adult clam during a generation) to obtain the progeny \((fN_i)\). The number of progeny was then multiplied by \(D_{ij}\) (see previous section) to determine their distribution among habitat patches.

Under the influence of migration (i.e., dispersal events) and in the absence of drift, the allele frequencies between pairs of sites were progressively homogenised. The aim of the model was thus to identify which pairs of sites were homogenised more quickly than others to test our hypothesis about dispersal and test the relative importance of geographic distance, oceanographic currents, giant clam abundance, and habitat patchiness.

**Comparison of simulated versus observed genetic structures**

The simulated genetic structures obtained from the simulation approach described above were then compared to observed genetic structure. Mantle biopsies of 849 *T. maxima* spread over 23 sampled sites were conducted from 2008 to 2014 (Fig 2, Table 2) and each individual was genotyped at 15 microsatellite loci as described in Grulois et al. [46].

Genetic diversity within samples was estimated using observed \((H_{obs})\) and Nei’s unbiased expected heterozygosity \((H_{exp})\) in GENETIX version 4.03 [47]. Single and multilocus \(F_{IS}\) were estimated using Weir & Cockerham’s [48] fixation index and deviations from Hardy-Weinberg equilibrium \((HWE)\) were tested using Fisher’s exact test using a Markov-chain randomization \((1000 \text{ dememorizations}, 100 \text{ batches}, \text{ and } 1000 \text{ iterations per batch})\) in GENEPOP version 3.4 [49] as implemented for online use (http://genepop.curtin.edu.au/). Significance levels for multiple comparisons of loci across samples were adjusted using a standard Bonferroni correction [50].

Since null alleles were detected in our samples, genetic divergences among samples were estimated in FREENA [51] (i) using the \(F_{ST}\) estimates of Weir & Cockerham [52] and the ENA method that provides unbiased \(F_{ST}\) estimates, and (ii) using a direct estimation of pairwise values for the Cavalli-Sforza & Edwards [53] genetic distance following the so-called INA method [51] that decreases the bias in genetic distance estimation observed when null alleles are present. Genotypic differentiation among samples was further tested with an exact test (Markov chain parameters: 1000 dememorizations, followed by 1000 batches of 1000 iterations per batch) with the original dataset, and the \(P\)-value of the log-likelihood \((G)\) based on the exact test [54] was estimated in GENEPOP version 3.4. Significance levels for multiple comparisons of loci across samples were adjusted using a sequential Bonferroni correction [50].

Among the dispersal kernels tested (IBD and IBOD), we identified those most similar to the observed genetic structure by comparing the observed and simulated standardized Cavalli-Sforza & Edwards’s [53] genetic distances. The simulated genetics distances \((d_s)\) were
compared with the observed genetic distances \( (d_o) \), using Mantel’s correlation coefficient for matrices.

**Sensitivity analyses of the genetic structure to seascape features**

Among all dispersal kernels tested in the previous sections, the one that simulated a genetic structure most similar to the observed structure was then used to evaluate the importance of reefscape composition and configuration in genetic patterns. The importance of reefscape composition was evaluated by degrading the high resolution habitat map (L3) to coarser habitat maps (down to L2 and L1). This provided three maps of habitats, each characterized by a more or less accurate spatial distribution of \( T_{\text{maxima}} \) density (Fig 3).

To evaluate if reefscape configuration (i.e. habitat continuity) and composition need to be taken into account for modelling genetic structure accurately, we compared the correlation between the observed and the simulated genetic structures obtained when considering the original habitat maps with the correlation obtained when degrading the original habitat map. Degradation of the original habitat map was performed by progressively reducing habitat area between populations surveyed for genetic data. For this, we respectively tested decreases from 20% to 80% of the initial habitat area. Patches that were removed were randomly chosen among patches not surveyed for genetic samples. These scenarios represented the process of habitat fragmentation that generates habitat loss and increases geographical distance between remaining habitat patches. All scenarios tested in this study are summarized in Table 4.

**Results**

**Giant clam observed genetic diversity and structure**

Raw microsatellite data can be found at https://figshare.com/s/a00f8be352aae916c147. Over all samples, the number of alleles per locus ranged from 11 to 40 (mean 26.8). The observed heterozygosities ranged from 0.493 to 0.610, and the expected heterozygosities ranged from 0.752 to 0.831 (See S1 Table). Significant deviations from HWE were observed in all samples, with multilocus estimates of \( F_{\text{IS}} \) ranging from 0.273 to 0.402, showing in all cases heterozygote deficiencies. However, six loci were in HWE in nearly all samples, suggesting that heterozygote deficiencies in the remaining nine loci were caused by the occurrence of null alleles (confirmed by the presence of null homozygotes at those nine loci).

The Efate sample (Vanuatu) was significantly differentiated from other sampled sites with pairwise Cavalli-Sforza & Edwards’ genetic distance values ranging from 0.388 to 0.467 (all \( P \)-values < 0.01, see S2 Table). The Chesterfield sample was also divergent from New Caledonia’s and Vanuatu’s samples (Cavalli-Sforza & Edwards’ genetic distance from 0.306 to 0.412, though not significant). Lower levels of genetic differentiation were observed among the sampled sites of New Caledonia (Cavalli-Sforza & Edwards’ genetic distance values from 0.235 to 0.38), but some sites in the north of New Caledonia (BEL and COO), the Loyalty islands (TIG) and the south of New Caledonia (GOR) held higher (but not significant) genetic distances with other sites. The KUA sample was also characterized by higher genetic distances with other sites (from 0.345 to 0.401), but was possibly biased by the low number of specimen collected at this station (\( n = 11 \), Table 2). This likely biased sample was excluded from further analyses.

**Dispersal kernel, connectivity matrices, and genetic structure**

The bio-physical oceanographic models showed that larvae released in the northern part of Vanuatu archipelago (above 17°S) tended to be exported northwestward by the North Vanuatu
Jet (NVJ) where no patch of suitable habitat exists. The occurrence of successful exports of larvae from Vanuatu to New-Caledonia progressively increased for release points further south, but successful exports were restricted to the Loyalty Islands, New Caledonia’s East coast, and Entrecasteaux reefs through the East Caledonian Current (ECC), the North Caledonian Jet (NCJ) and the South Caledonian Jet (SCJ). By contrast, a significant fraction of larvae could cross the distance between the north part of New Caledonia and the Chesterfield Islands using eddies issued from the NCJ and the Sub-Tropical Counter Current (STCC). This provided a connectivity matrix with higher connections among habitat patches of New Caledonia than between Vanuatu, New Caledonia, and Chesterfield (Fig 4B).

The dispersal kernel that provided the best match with genetic data was the IBD scenario with $a = 0.95$ in Eq 1 (S2 Fig). The connectivity matrix for this scenario is provided in Fig 4A and suggests high self-recruitment and restricted dispersal. For this scenario, dispersal was restricted to very proximate patches (e.g. between AST and BB or between LIF and TIG), and never occurred beyond 150 km (S3 Fig).

When considering an IBD pattern for dispersal, the gene flow model adequately isolated Vanuatu and Chesterfield from New-Caledonia, but did not adequately mimic the genetic structure among New-Caledonia’s reefs (Fig 5). By contrast, the IBOD model failed to project isolation for Chesterfield islands, and estimated higher relative isolation for the south group than was observed in the empirical genetic data (Fig 5).

**Sensitivity analyses of genetic structure to seascape features**

Degrading the original habitat map by reducing habitat area significantly decreased the correlation between simulated and observed genetic distances ($F = 13.8, p < 0.01; $Fig 6$)$. When only 20% of the initial habitat area was maintained (i.e. 80% fragmentation level), correlation decreased from $0.66 \pm 0.08$ to $0.50 \pm 0.14$ for the L3 scenario, from $0.58 \pm 0.14$ to $0.47 \pm 0.18$ for the L2 scenario, and from $0.63 \pm 0.07$ to $0.44 \pm 0.22$ for the L1 scenario. The influence of habitat composition was also significant ($F = 11.6, p < 0.01$), with correlation values lower for the “L1” and “L2” scenarios than for the “L3” scenario, especially when habitat fragmentation was 40%, 60% and 80%.

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Table 4. Summary of all scenarios tested in this study and the corresponding correlations between the simulated and observed genetic structure.

<table>
<thead>
<tr>
<th>Scenarios</th>
<th>Dispersal kernel</th>
<th>Habitat fragmentation</th>
<th>Habitat composition</th>
<th>Mantel coefficient of correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
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<td>0</td>
<td>L3</td>
<td>$0.66 \pm 0.08$</td>
</tr>
<tr>
<td>2</td>
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<td>20</td>
<td>L3</td>
<td>$0.58 \pm 0.12$</td>
</tr>
<tr>
<td>3</td>
<td>IBD</td>
<td>40</td>
<td>L3</td>
<td>$0.56 \pm 0.12$</td>
</tr>
<tr>
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<td>IBD</td>
<td>60</td>
<td>L3</td>
<td>$0.53 \pm 0.14$</td>
</tr>
<tr>
<td>5</td>
<td>IBD</td>
<td>80</td>
<td>L3</td>
<td>$0.50 \pm 0.14$</td>
</tr>
<tr>
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<td>IBD</td>
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<td>L2</td>
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</tr>
<tr>
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<td>$0.50 \pm 0.19$</td>
</tr>
<tr>
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</tr>
<tr>
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<td>L2</td>
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</tr>
<tr>
<td>10</td>
<td>IBD</td>
<td>80</td>
<td>L2</td>
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</tr>
<tr>
<td>11</td>
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<td>L1</td>
<td>$0.63 \pm 0.07$</td>
</tr>
<tr>
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<td>IBD</td>
<td>20</td>
<td>L1</td>
<td>$0.51 \pm 0.14$</td>
</tr>
<tr>
<td>13</td>
<td>IBD</td>
<td>40</td>
<td>L1</td>
<td>$0.40 \pm 0.20$</td>
</tr>
<tr>
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<td>IBD</td>
<td>60</td>
<td>L1</td>
<td>$0.41 \pm 0.16$</td>
</tr>
<tr>
<td>15</td>
<td>IBD</td>
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<td>L1</td>
<td>$0.44 \pm 0.22$</td>
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https://doi.org/10.1371/journal.pone.0178239.t004
Discussion

In this study, we simulated the expected genetic differentiations of giant clams across the study area given the reefscape composition and configuration and under various scenarios of larval dispersal. The simulated genetic structures were compared with the observed genetic structure established from empirical genetic data. This general framework is an example of the multidisciplinary approach recommended by Selkoe et al. [12]. The modelling approach here involves (i) maps of habitat suitability of the species over the study area; (ii) biophysical modelling for propagule dispersal between all habitat patches; (iii) gene flow modelling that accounted for multi-generational stepping stone processes; and (iv) sensitivity analyses to evaluate the extent to which reefscape configuration and composition are necessary for modelling
This simulation approach led to similar conclusions than a regression approach (see S2 File), but could explicitly test the influence of environmental features on genetic patterns, while taking into account many of the complex processes that drive larval connectivity. This approach is a promising alternative to the simple regressions classically used in landscape or seascape studies, when habitat fragmentation/continuity cannot be resumed.

**Fig 5.** Comparison between observed and simulated genetic structures for *Tridacna maxima*. A) Observed genetic distances from empirical data, obtained from 15 microsatellite loci. B) Genetic distances simulated by the isolation by distance (IBD) model. C) Genetic distances simulated by the isolation by oceanographic distance (IBOD) model. Genetic distances are Cavalli-Sforza & Edwards’s genetic distances [53], normalized to have maximum of 1.

https://doi.org/10.1371/journal.pone.0178239.g005

**Fig 6.** Correlations between observed and simulated genetic structures for the various scenarios of landscape composition and configuration. Landscape composition was determined by the three levels of the reef typology described in Table 2. Landscape configuration is determined by habitat fragmentation (0% refers to the initial habitat maps, while 20%, 40%, 60% and 80% refer to reduction levels of habitat area compared to the initial habitat map). The coefficient of correlation is the Mantel coefficient of matrix correlation. We used the dispersal kernel that provided the best congruence between the observed and simulated genetic distances (IBD). Bars are medians and error bars are quantile 95%.

https://doi.org/10.1371/journal.pone.0178239.g006
by a single variable that would influence genetic patterns in a linear way, and interact with other factors (e.g., oceanography) also in a linear way.

The approach adopted here has a number of limitations that are important to acknowledge. First, except between Vanuatu and other samples, genetic differentiation levels were found to be low in the study area. Landscape/seascape genetics traditionally aims to evaluate genetic structure and relate it to environmental features. Faint genetic structure requires addressing the problem in a different way: which environmental features can explain the significant or non-significant genetic differentiations observed among sites? While lack of structure is not usually the focus of landscape geneticists [13], it is likely to be an important focus for seascape genetics because marine species usually have low levels of genetic structure at the spatial scale of concern for management purposes [55, 56, 57]. The limited (but present) genetic structure observed for giant clams in our study (see Vanuatu, S2 Table, Fig 5) is therefore representative of, and a useful example for, future studies in the seascape literature.

Second, the multi-disciplinary approach performed here required extensive datasets that were not easy to collect and bring together. Habitat maps were readily available, but a typology adapted to the current knowledge on *T. maxima* density in the literature needed to be established (Table 3) to represent *T. maxima* habitats adequately [31]. For example, comprehensive data on giant clam density in each habitat patch were not available at the wide geographical scale of our study. Specifically, we had to use the density of giant clams in each habitat type to calculate giant clam abundances for each habitat patch. We used a dataset from the literature that itself involved a substantial sampling effort (n = 31 sites surveyed over 6 shallow geomorphological units), though many regions and species do not have such data readily available. Despite the effort, we still lacked fine density estimations for reefs of variable depths (Table 3). Our results nevertheless demonstrated that more accurate characterization of reefscape features enhanced our understanding of genetic structure.

Third, the model used in this study projected gene flow on the basis of migration only (through larval dispersal). Other evolutionary forces (selection, mutation, and genetic drift) were neglected, but their influence on the results should be assessed in future research programs. Mutation is likely unimportant at the small spatial and temporal scales that are addressed in this study. Selection may affect parts of the genome including neutral genomic regions [58], but it was unlikely the case in our study because all microsatellite loci provided similar Fst values (S3 File). Then, genetic drift may accentuate the divergence of low density (i.e. low population size) clam habitat. Integrating processes involved in demographic connectivity and genetic connectivity is not an easy task and requires careful consideration of the life history and natural history of the species being studied. For long-lived species like giant clams, special consideration will be required in future research to handle overlapping generations in the model. Overlapping generations may modify gene flow and the relative influence of evolutionary forces on genetic structures [59].

Finally, we used the Mantel correlation coefficient to compare simulated *versus* observed genetic distances, but the Mantel test has been recently criticized due to low robustness to spatial autocorrelation [60, 61]. While using the Mantel correlation may have underestimated correlation coefficients between simulated *versus* observed genetic distances, it is unlikely to have changed the conclusions of our study which are based on a relative comparison of correlation coefficients between scenarios and not on the absolute value of correlation.

**The relatively poor performance of IBOD compared to IBD in our study**

Only by considering IBD with limited dispersal ($a = 0.95$ in Eq 1) did the model project higher isolation for Vanuatu and Chesterfield, in agreement with observed genetic data (Fig 5). This
result suggested that larval dispersal was restricted to short distances, usually several tens of kilometres, and rarely exceeded 150 km for giant clams in the New Caledonia and Vanuatu area (S3 Fig), a distance supported by an independent analysis which estimated the genetic patch size from a spatial auto-correlation analysis from the New Caledonia genetic survey (see S4 File). This result is also in agreement with dispersal distances estimated in other areas for species with relatively short PLD [62, 63, 64].

The IBD model projected relative isolation for Vanuatu and Chesterfield that was similar to the observed genetic divergence (Fig 5), but this was not the case for IBOD. Conversely, none of the tested scenarios could explain the genetic structure observed at finer scales, such as the higher (but not significant) isolation found for some sites in the north of New Caledonia (BEL and COO), the Loyalty islands (TIG) and the south of New Caledonia (GOR). The relatively poor performance of IBOD model compared to IBD may result from the fact that we only considered ocean currents in IBOD, while neglecting connections through lagoon waters. Much of *T. maxima* specimen in the area are found in the lagoon of New Caledonia, and for them the current approach does not accurately model larval dispersal at the beginning and end of the dispersal period. The movements of lagoon waters are generally driven by tide, wind and swell [65] and reef configuration [66]. In New Caledonia wide and geomorphologically complex lagoon [30], current intensities and directions remain poorly characterized especially around shallow reefs, but may produce a range of possible distances for larval dispersal [67] which may shape the fine scale genetic structure over the long run.

Our biophysical model (IBOD) considered the PLD, larval survival, and ability for larvae to settle when crossing a suitable substrate. Available computer time limited our ability to run dedicated sensitivity analyses for each of these parameters, but all are expected to influence the dispersal kernel [40] and warrant further investigation. A number of other processes also deserve more attention for future modelling of larval dispersal, including 1) the movement of gametes before fertilization, which last several to tens of hours for giant clams [40] and may significantly influence dispersal kernel; 2) larval swimming speed, which may also affect the path of larvae and is low for *T. squamosa* (104.0 to 1010.6 μm.s\(^{-1}\), [40]) but remains unestimated for *T. maxima* (though larval behaviour experiments reveal active swimming, [39]); 3) factors inducing settlement, including crustose coralline algae [40] or conspecifics [39]; and 4) spatial differences in larval survival, such as from spatial variation in predation or food availability. Finally, it is worth noting that despite the recent advances in characterizing the circulation around New Caledonia [32], ROMS models continue to have difficulty simulating currents along the coastlines, particularly in the Loyalty channel where the modeled Vauban current is too strong. Accurate knowledge of eddy dynamics is also lacking for the West and South part of New Caledonia. This could bias dispersal kernels in these areas and explain the low performance of IBOD models. However, the model available for New Caledonia remains far better than what is available in most coral reef ecosystems and has been validated extensively against empirical observations (see S1 File). Since the IBOD model was not supported by genetic data, it was not used in our analysis to test the hypothesis that reefscape configuration and composition influence genetic structure. Future research needs to improve the accuracy of the oceanographic model so that a reliable evaluation of the effect of reefscape features in an IBOD model can be performed.

Reefscape genetics: Accounting for reef distribution in seascape genetics

The idea that habitat fragmentation and composition can impact demographic and genetic connectivity patterns has strong foundations in the terrestrial literature [13, 6]. The marine realm,
by contrast, has often been considered a relatively continuous environment devoid of barriers for dispersal. The concept of “seascape genetics” only recently emerged in marine ecology as studies began linking connectivity patterns for geographic features. Thus far, seascape genetics has mostly used meso-scale oceanic features (i.e., currents, [68]) to describe this seascape, even for species and populations that spend most of their life in coastal benthic habitats. Specifically in tropical marine ecosystems, the diversity of reef configurations induced by contrasting patterns of reef patchiness and other reefscape features (e.g., reef geomorphology, lagoon enclosure) may also shape connectivity [11]. In this study, we highlight that even relatively simple environmental features inherent from the landscape ecology literature can be important for modelling and understanding genetic patterns for a marine species. These environmental features, related to hard bottom habitat composition and configuration, can therefore enhance our understanding of genetic patterns in coral reef ecosystems. To accurately model the genetic structure of reef species in a region, we stress the need for a specific approach that integrates reef habitat composition and configuration with oceanic and lagoon currents, hence a “reefscape genetics” approach.

Reefscape features as proxy for connectivity: Perspectives for future research

Population connectivity often enhances resilience to climate change and other anthropogenic impacts on ecosystems [69]. Loss of connectivity due to habitat destruction and global warming is therefore raising concerns. Connectivity has been the focus of many investigations, but to date, seascape genetic studies typically focus on larval transport (e.g., oceanographic currents) rather than production and settlement sites. Accurately modelling larval transport is often challenging and costly, however, and requires specialized computing capabilities. To date, most coral reef ecosystems lack such detailed information, which limits the ability of biophysical models to help characterize demographic and genetic connectivity in a wide variety of places. In this context, reef biologists and managers would benefit from simple and accurate predictors of gene flow. We propose that reefscape genetics and a new focus on habitat geography can offer a fresh perspective.

Seascape features can provide useful spatial surrogates for population genetic structure that are potentially relevant for a wide range of marine organisms. Our study concluded that habitat composition and configuration significantly influenced genetic patterns of giant clams. Further investigation is needed to assess the robustness of this result to species traits (e.g., PLD), but habitat maps and other reefscape features may provide a new path for marine ecology in regions where more detailed information on ocean currents are lacking.

We suggest future research to investigate the following directions. First, one important advance will be to quantify the relative influence of a wider variety of reefscape features on genetic structure. We here demonstrated that reefscape composition and configuration can enhance our understanding of population connectivity through a reduction of habitat area and a degradation of habitat map, but the influence of reefscape features like reef enclosure (e.g., atoll versus island reef) should also be investigated. It would also be helpful to compare results from reefscape genetics among taxa and identify proxies of connectivity robust enough to be related to particular species traits, including PLD. Over the long run, this could help elevate reefscape genetics from a population to community scale and help characterize the link between reefscape features and biodiversity broadly. Finally, it will be useful to characterize how reef use, reef degradation, and other aspects of environmental change affect genetic connectivity, and to assess the consequences for reef management and for the identification of priority areas for conservation. These three recommendations are not exhaustive,
but would help reinforce reefscape genetics as a new path for marine ecology and conservation science.

Supporting information

S1 Table. Summary of genetic diversity at 15 microsatellite loci from *Tridacna maxima* samples. n: number of sampled individuals; \( H_{\text{obs}} \): observed heterozygosity; \( H_{\text{exp}} \): Nei’s unbiased expected heterozygosity; \( F_{IS} \): Weir and Cockerham’s (1984) estimate of Wright’s (1951) fixation index (italic type indicate significant deviations from HWE after standard Bonferroni correction). Full names of abbreviated sampled locations are given in Fig 2.

(PDF)

S2 Table. Observed pairwise genetic differentiation among *T. maxima* sampled locations. Above diagonal: pairwise \( F_{ST} \); below diagonal: Cavalli-Sforza and Edwards (1967) genetic distance used for all comparisons with simulated genetic structures. Significant genotypic differentiations are indicated in italic for \( P \)-value < 0.01 and in bold for significant after sequential Bonferroni correction. Full names of abbreviated sampled locations are given in Fig 2.

(PDF)

S1 Fig. Relative probability of survival for *Tridacna maxima* larvae over the competency period. Competency period considered is from 9 to 19 days, and the relative probability of survival was considered for calculation of dispersal probability in the IBOD model.

(PDF)

S2 Fig. Performance of IBD models to reproduce the observed genetic structure as a function of parameter \( a \) (see Eq 1). A) Results obtained when using the mantel coefficient of matrix correlation. B) Results obtained when using the linear R-squared. Points are medians and error bars are quantile 5% and 95%.

(PDF)

S3 Fig. IBD dispersal kernel that provided best congruence between the simulated and observed genetic structures for *T. maxima* in the New Caledonia and Vanuatu area.

(PDF)

S1 File. Background surface circulation and model configuration.

(PDF)

S2 File. Linear regression models between genetic distance, geographic distance, oceanographic distance, and habitat continuity.

(PDF)

S3 File. \( F_{ST} \) values per locus obtained with and without using ENA.

(PDF)


(PDF)

Acknowledgments

We are grateful to the R Core Team for the design and freely available software R3.1.0, and to A. Renaud, P. Dumas and J. Butscher for their help in sampling giant clams. Thanks to Firas Al-khalil for his advice concerning the pathfinding algorithm. This is ENTROPIE contribution #213.
Author Contributions
Conceptualization: SVW SA.
Data curation: CF JT DG.
Formal analysis: SVW JT DG CF SA JL.
Funding acquisition: CF NGM.
Investigation: SVW.
Methodology: SVW MLP.
Project administration: SA CF.
Resources: CF SA.
Software: SVW.
Supervision: SA NGM.
Validation: SVW.
Visualization: SVW.
Writing – original draft: SVW.
Writing – review & editing: SA MLP CF JL JT DG SVW NGM.

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