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Fish community structure in relation to environmental variation in coastal volcanic habitats

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A B S T R A C T

This study constitutes one of the few works conducted on the colonization of recent lava flows by fish communities in coral reef environment. It attempted to describe the mechanisms of colonization and environmental variables influencing the structure of communities on the lava flows of the Piton de la Fournaise volcano at Reunion Island. It also proposed some hypotheses on the mechanisms of selection and speciation of endemic species of the southwestern Indian Ocean region, particularly diversified on the most recent lava flows of the Piton de la Fournaise, and provided a model of distribution of environmental and fish assemblage variables in four homogeneous patterns of colonization (landscapes) retained within the study area. The results showed fish community structure strongly related to measured environmental variables, of which depth, width of the island shelf, age of the lava flow and distance to the most recent lava flow were the most influential. It also appeared that the maximum endemic species richness was measured on the shallower stations, with narrow insular shelf, located on recent lava flows, near the lava flow from the eruption of 2007 (latest volcanic event). These characteristics define a harsh environment, to which endemic species seem well-adapted due in particular to their strong demographic and dietary flexibility. Finally, the patterns of colonization were characterized by significantly different environmental variables. The four homogeneous areas, defined by two geomorphological features, basalt blocks and drop-offs, nested within two locations, inside and outside the volcanic enclosure (current caldera), hosted fish communities whose dominant diets mainly depend on the nature, the abundance and the accessibility of available food resources. Overall, the study highlights that during early stages of ecological succession, site selection by fish communities is highly dependent on environmental constraints, especially of volcanic origin, but the physical habitat selection (geomorphology, substrate rugosity, percentage of soft substrate) seems to have little importance, while at least on the trophic structure and the distribution of endemic species.

1. Introduction

The ecological requirements of tropical marine fishes have been extensively studied for 50 years including their community structure and interactions of communities with environmental variables such as substrate rugosity (e.g. Chabanet et al., 1997; Gratwicke and Speight, 2005), benthic organism cover (e.g. Bell and Galzin, 1984; Chabanet and Faure, 1994), depth (e.g. Booth and Wellington, 1998; Friedlander and Parrish, 1998), proportion of soft substrate (e.g. Khalaf and Kochzius, 2002; Travers et al., 2010), and intensity of a volcanic disturbance (e.g. Godwin and Kosaki, 1989; Pinault et al., 2013b). However, no model attempting to explain fish community structure in relation to environmental variables has been constructed for an active volcanic area.

Otherwise, the southwestern Indian Ocean (SWIO) region, comprising a main landmass, Madagascar, and several island archipelagos such as Comoros, Mascarenes and Seychelles, each with different origins and ages, hosts a high proportion of endemics and is highly threatened by human activities, hence its classification as a marine biodiversity hotspot (Bellard et al., 2013; Roberts et al., 2002). This region constitutes a natural laboratory for studying diversification processes in coastal marine taxa, whose rocky shore species have been yet little studied (Postaire et al., 2014).

This lack of information can be attributed to the traditional focus on coral reefs, often considered to host the highest biodiversity in the tropics (Sale, 1991), and to difficulties in standardization of sampling...
methods to the geomorphological and ecological specificities of rocky shore habitats, including volcanic areas (Pinault et al., 2013b).

Reunion Island is one of the few places where volcanic activity is frequent, with 27 eruptions occurring between 1998 and 2007 and a mean frequency of an eruptive phase every 9 months over the past century (Tanguy et al., 2011). It is also one of the few places where lava frequently flows into the ocean (Michon and Saint-Ange, 2008). Community development on lava flows is of considerable interest to the natural history of Reunion Island and the Mascarene Archipelago. Although terrestrial succession on lava has been well studied at Reunion (Strasberg, 1995), the development of fish assemblages has been monitored only by Pinault et al. (2013b, in press) on the submerged flows of the Piton de la Fournaise.

The aims of this study were to (1) estimate the relative influence of environmental variables on the fish community structure settled on recently formed habitats regularly subjected to acute disturbances, (2) explain the distribution of endemic species in the SWIO rocky shore habitats exploring their adaptation to environmental variability, and (3) propose a fish colonization pattern on the surveyed volcanic area according to the main underwater landscapes and their specific environmental variables.

2. Material and methods

2.1. Study site

Reunion is an island of the Mascarene Archipelago, with Mauritius and Rodrigues Islands, 690 km east of Madagascar (21.06°N, 55.33°E) (Fig. 1). Of recent volcanic origin (between 1.5 and 8 Ma), the three islands have undergone alterations following successive eruptive phases. Reunion Island consists of two volcanoes, the Piton des Neiges and the Piton de la Fournaise: The former has been extinct for c. 70,000 years, while the latter is still active with effusive eruptions (Chevallier and Vatin-Perignon, 1982).

The coastline of Reunion is characterized by a narrow insular shelf and steep slopes. The west-southwest coast is urbanized, with fringing coral reefs extending for about 25 km (12% of the total coastline) (Pinault et al., 2013a). The north coast, relatively shielded from the effects of trade winds, is also densely populated. It receives 9 of the 13 perennial river mouths and is characterized by turbid waters and a high proportion of soft substrate. The southeast region of the island, deeply marked by recent and ongoing volcanic activity of the Piton de la Fournaise, is sparsely inhabited. Most of the historic eruptions of the Piton de la Fournaise (2002, 2004, 2005, 2007) have been from vents located inside the walls of the current caldera or volcanic enclosure (VE), at various elevations along the rift zone, but three of the most recent four large-volume deep-seated eruptions (1977, 1986, 1998) have had at least one vent located outside of the caldera rim, along an extension of the intracaldera rift zone (Vigouroux et al., 2009). This volcanic area is exposed to trade winds and is characterized by regular swells, sometimes strong, particularly during the austral winter (Bollard et al., 2013), and by widespread runoff and percolation generated by heavy rainfall.

The shoreline of the VE, which forms the southern area of the study site, consists of sea cliffs 3 to 8 m high that continue vertically underwater to depths of 3 to 5 m. The deeper areas of the flows consist largely of unconsolidated lava boulder, loose rocks, and rubble, with the bottom sloping into deep water at an angle of c. 30°. The area immediately to the north of the VE, which forms the central area of the study site, is on a prehistoric flow of undetermined age. First evidence of an eruption of the Piton de la Fournaise dates from 1640, but many flows occurred before that date (McDougall, 1971). The northern area of the study site consists of the 1977 flow, which entered the sea outside the VE, creating a shoreline c. 750 m wide with sea cliffs 3 to 5 m high and narrow black sand beaches. The substratum immediately offshore, deposited by prehistoric and the 1977 flows, consists of 1 to 3 m diameter angular boulders and slopes steeply to a depth of 8 m. Between 8 and 20 m, the bottom slopes downward more gradually; beyond the 20 m contour the bottom once again drops away at a steep angle. The substratum between 8 and 20 m consists of compact wave-smoothed lava with high algal cover. Pocillopora verrucosa (Ellis and Solander, 1786), Pocillopora eydouxi Milne-Edwards and Haime, 1860, Pocillopora meandrina (Linnaeus, 1758), and Pocillopora meandrina Dana, 1846 are the most abundant corals, but cover is sparse and tends to be concentrated in the northern reaches of the area. Well-developed coral communities were found between 10 and 25 m north of the study site on the 1977 flow and at the Caesari Rock and Waterfalls Bay areas (Fig. 1, transects T5, T6, T7, T8, T9, T10).
1–3). Although the health status of the coral communities was good in the two northernmost areas (transects 1–2), the corals were partially dead or necrotic in Waterfalls Bay (transect 3).

2.2. Sampling method

Fieldwork took place over an eight-day period in late November 2011 (ustral summer). Surveys were taken at thirty-eight–eight stations at depths from 5 to 30 m within 10 linear transects perpendicular to the shore (Fig. 1). Each transect contained four sampling stations (5–10, 11–15, 16–20, and 21–30 m) except for T2 and T10, which contained three stations (10–15, 16–20, and 21–30 m) because high surge conditions and limited visibility due to a “mist” of fine air bubbles made it impossible to survey the shallow sub-tidal communities on the final survey day. The transects were located along the shore in two sectors inside and outside the VE, on flows of different ages. All surveys took place between 0900 and 1600 h.

The deepest end of each transect (30 m) was located by dropping a weighted float overboard after recording the station position with a Garmin eTrex GPS receiver (www.garmin.com). Two trained divers equipped with slates, pencils and data sheets moved randomly over an estimated area of 300 m² at depths of 21 to 30 m, recording each observed species using underwater visual censuses (UVC) techniques, and estimating hard substrate rugosity, soft substrate percentage, coral and algal cover using a habitat assessment score following the methods of Pinault et al. (2013a) (station 1). When 15 min had elapsed, the divers moved toward shore and repeated the protocol at 16 to 20 m depth (station 2), then 11 to 15 m (station 3), and finally 5 to 10 m (station 4). The UVC is a non-destructive, capture-independent method, superior to collection techniques, especially in sensitive areas. Underwater visual censuses require little post-processing, whereas collections take many more hours to complete. The method does, however, tend to underestimate the numbers of small, nocturnal, and cryptic fish (Ackerman et al., 2004; Fowler, 1987; Harmelin-Vivien et al., 1985; Pinault, 2013; Willis, 2001).

In order to minimize this bias, a stratified sampling method consisting of dividing the population into homogeneous sub-groups can be implemented (Labrosse et al., 2002). In this study, two strata were created, based mainly on the behavior of the observed fish species: (1) demersal and pelagic species such as Labridae, Pomacentridae, and Carangidae and (2) mimic, burrowing and cave species such as Gobiidae, Scorpaenidae, and Muraenidae. Each diver surveyed one stratum in an identical manner (same sampling time, way and surface).

2.3. Fish assemblage variables

Recorded species were assigned to eight groups according to diet, based on extensive published data (Froese and Pauly, 2012; Hiatt and Strasburg, 1960; Hobson, 1974; Myers, 1999): (1) herbivores, grazing on microalgae or macrophytes; (2) omnivores, feeding opportunistically on available food, including organic debris; (3) browsers of sessile invertebrates, feeding mainly on coral polyps, but also on other fixed organisms; (4) diurnal and (5) nocturnal carnivores, consuming benthic invertebrates; (6) piscivores, preying on other fish; and (7) diurnal and (8) nocturnal planktivores, capturing plankton, mainly animal. Only the primary trophic assignment of each species was included in the analyses. When authorities presented conflicting information for a given species, the most common trophic assignment was selected.

Ecological traits were defined based on species biological and behavioral characteristics such as asymptotic length (L∞), minimum and maximum depths, migratory behavior and fecundity. The asymptotic length was provided by the von Bertalanffy growth equation parameters (Bertalanffy, 1938), available in Froese and Pauly (2012). The minimum and maximum depths were found in Lieske and Myers (1994). The migratory behavior was determined using McPherson and Jetz (2007). Species closely associated with their habitat and generally characterized by a narrow species range-size were classified as sedentary. Conversely, seasonal visitors and species exhibiting local movements within the study area were treated as nomadic. The species fecundity was deduced from Froese and Pauly (2012). Species with minimum population doubling time less than 15 months were classified as highly fecund. Others were indistinctly considered as moderately or least fecund.

Endemic species of the SWIO region (SWIO species), in the sense of Pinault et al. (2013b), were considered as the most representative of the patrimonial value of the stations. They were treated separately from the non-endemic species (non-SWIO species) in proper analyses. A biogeographic pattern was provided per species by the online system AquaMaps (Kaschner et al., 2010), a tool using a niche modeling approach (Hutchinson, 1957) to generate predicted distributions of marine species based on available distribution data or described ecological tolerances. In essence, provided that a minimum acceptable level of information is available, the tool will generate a predicted distribution for any species, calculated by the probability of occurrence in the entire ocean. Predicted distributions, based on ecological tolerances, often present ranges that exceed the actual known distribution of species, because the geographic distributions of suitable ecological variables usually extend over a much greater area than the known ranges of species (Rees, 2008). This theoretical approach was essential in order to assess the distribution of rare species and especially species not previously recorded from Reunion Island. This method, however, ignores the evolutionary constraints, providing ranges that would be possible if dispersal was not limited by geological history, which is particularly complex in volcanic oceanic islands (Postaire et al., 2014). It is therefore possible that some model-based distribution patterns are broader than what exist in reality. Some species identified as restricted to the SWIO area could be thus restricted to a smaller range, as the Mascarene Islands.

2.4. Additional environmental variables

The width of the island shelf in meters and the distance from the most recent lava flow (2007) in kilometers were estimated by geographic information system (GIS), using Mapinfo professional software (www.mapinfo.com). For each station, the width of the insular shelf was measured transversely to the shore from the high tide line to the −100 m isobath using the bathymetric map of the naval hydrographic and oceanographic service (SHOM).

The age of the lava flows was calculated in years before the survey period. The flow from the 2007 eruption was, for example, estimated at 4 years. The four transects with undetermined ages (Fig. 1, T2–5) were considered to have more than 371 years, the first evidence of an eruption of the Piton de la Fournaise dating back to 1640 (McDougall, 1971).

2.5. Underwater landscape features

Despite the diversity of underwater landscapes observed during survey, only two geomorphologies: basalt blocks and drop-offs and two locations: outside (Fig. 1, T1–5) and inside VE (T6–10) were retained. The nested design of these two factors led to the creation of four main underwater landscapes: basalt blocks outside VE, basalt blocks inside VE, drop offs outside VE, and drop offs inside VE.

Basalt blocks consist of large lava boulders and loose rocks caused by the erosion of the sea cliffs that constitutes the coastline of the study site. These blocks accumulate over wave-smoothed compact lava from successive eruptive phases, sloping into deeper water at a slight angle. Drop-offs consist of compact lava cliffs, sloping steeply on sedimentary patches including a mixture of olivine sand and centimetric scoria gravel. The basalt cliffs frequently have columnar appearance typically known as basalt organ pipes, because of their regular hexagonal shape.

The area outside VE consists of wave-smoothed volcanic rocks, generally more colonized by sessile marine fauna and flora than the area inside VE. Littoral benthic ecosystems are inhabited by a multitude
of herbivorous and lithophagous organisms that strongly influence the underwater landscape mosaic by their selective grazing and bio-
erosive action. In contrast, the area inside VE, including the most recent lava flows of the Piton de la Fournaise, exhibits mineral landscapes,
which consist of unconsolidated lava boulders, scoria gravel and rubble, with very steep slopes. The background colors are dark in crystal clear seawater.

2.6. Data analysis

To assess spatial variation, environmental variable values were com-
pared among underwater landscapes using a two-stage nested design in
analysis of variance (nested ANOVA, geomorphology nested within lo-
dation) (Montgomery, 1976). The normality of the statistical distribu-
tion of the environmental variables was assessed by a Shapiro–Wilk
test (Shapiro and Wilk, 1965). Bartlett’s test for homogeneity of vari-
ances (Snedecor and Cochran, 1989) was used to verify whether values
measured in different geomorphologies and locations had equal vari-
ances. When necessary, the one-parameter Box–Cox transformation
(Box and Cox, 1964) was implemented to reduce anomalies such as
non-additivity, non-normality, and heteroscedasticity (Saikia, 1992).

The relationships between fish assemblage structure (species occur-
rences) and environmental variables were elucidated using a canonical
correspondence analysis (CCA) (ter Braak, 1986). It is a fact that rare
species are often positioned as outliers in correspondence analysis ordi-
nations (Greenacre, 2013). Furthermore, statistical estimation problems
are posed by multicollinearity (ter Braak, 1986). In this sense, species
occurring at less than 10 stations and significantly correlated environ-
mental variables (correlation test — Pearson, 1957) were omitted from
analysis prior to conducting CCA. A Monte-Carlo randomization
test (1000 permutations) (McCune and Grace, 2002) was used to assess
the probability of the observed pattern being due to chance.

Multiple regression analyses (Efroymson, 1960) were then imple-
mented to test the relationships between fish assemblage variables
(diets and biogeographic patterns) and environmental variables. One
analysis was conducted per fish assemblage variable, integrating envi-
ronmental variables as predictors. For the same reasons as for the CCA,
significantly correlated environmental variables were excluded from
data prior to conducting multiple regression analyses.

Finally, it might be argued that different landscapes should exhibit
inherently different fish assemblage variable values, and that any ob-
erved among-site differences are due to geomorphology or location
rather than rugosity, depth, or other quantitative environmental vari-
ables. This hypothesis was tested only for the SWIO species richness,
rather than rugosity, depth, or other quantitative environmental vari-
able environmental variables as predictors. For the same reasons as for the CCA,
the probability of the observed pattern being due to chance.

3. Results

3.1. Spatial variation

The basalt blocks were characterized by significantly shallower
depths (nested ANOVA, p < 0.001), higher percentages of coral cover
(p = 0.016), and lower proportions of soft substrate (p = 0.028) than the
drop-offs (Table 1). The area outside VE was distinguished by signif-
icantly longer distances to the most recent lava flow (p < 0.001), older
ages (p = 0.001), higher percentages of coral cover (p = 0.001), and a
wider insular shelf (p < 0.001) than the area inside VE. Only the per-
centages of coral cover showed significantly different values between
g eomorphologies and locations. Thus, corals seemed to grow well on
basalt blocks outside VE. In contrast, algal cover and rugosity showed
no significant differences among underwater landscapes (p > 0.05).

3.2. CCA analysis

Width of the insular shelf, age of the lava flow and distance to the most
recent lava flow were highly correlated (Pearson test, p < 0.001) signifi-
cing that recent lava flows tend to increase the slope between the shoreline
and the depth at 100 m. Similarly, coral cover was positively correlated
distance to the most recent lava flow (p < 0.001) and negatively cor-
related with soft substrate (p < 0.001). The negative influence of the most
recent lava flow on coral communities may be the result of the stress suf-
fered by the entire coastal ecosystem within the study area at the latest
eruption of the Piton de la Fournaise in 2007. Finally, rugosity was nega-
tively correlated with algal cover (p < 0.001); thus, algal communities
seem to develop preferentially on wave-smoothed volcanic rocks. Since
these relationships caused multicollinearity problems, age of the lava
flow, distance to the most recent lava flow, and algal and coral covers
were excluded from further analyses.

The CCA revealed that spatial (Fig. 2A) and fish assemblage (Fig. 2B)
patterns are strongly related to the underwater landscapes of the
surveyed area.

The first axis, which is mainly a linear combination of the percentages
of soft substrate and the width of the insular shelf, separates stations out-
side VE (right of the axis) and those located inside VE (left of the axis).
Fish species that occurred mainly outside VE were characterized by higher
proportions of browsers of sessile invertebrates [14% vs. 9% inside VE,
Chaetodon guttatissimus Bennett, 1833, Forcipiger flavissimus Jordan
and McGregor, 1898, Pomacanthus imperator (Bloch, 1787)] , while those
occurring mainly inside VE presented greater percentages of omni-
vores [29% vs. 19% outside VE, Acanthus dussumieri Valenciennes,
1835, Acanthus mata (Cuvier, 1829), Chaetodon kleinii Bloch, 1790,
Pomachromis richardsoni (Snyder, 1909), Pseudanthias squamipinnis
(Peters, 1855)] and SWIO species [19% vs. 9% outside VE, Bodianus
macrourus (Lacepède, 1801), Cirrhichthys guichenoti (Sauvage, 1880),

Table 1

<table>
<thead>
<tr>
<th>Environmental variables (units)</th>
<th>Location</th>
<th>Geomorphology</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Outside volcanic enclosure n = 19</td>
<td>Inside volcanic enclosure n = 19</td>
</tr>
<tr>
<td>Dist. to the latest lava flow (km)</td>
<td>9.7 ± 0.8*</td>
<td>2.3 ± 2.9*</td>
</tr>
<tr>
<td>Age of lava flows (years)</td>
<td>323 ± 121*</td>
<td>7 ± 2*</td>
</tr>
<tr>
<td>Depth (m)</td>
<td>169 ± 5.3</td>
<td>15.0 ± 7.5</td>
</tr>
<tr>
<td>Coral cover (%)</td>
<td>19.8 ± 20.5*</td>
<td>3.9 ± 2.9*</td>
</tr>
<tr>
<td>Algal cover (%)</td>
<td>52.5 ± 13.9</td>
<td>45.2 ± 20.7</td>
</tr>
<tr>
<td>Soft substrate (%)</td>
<td>11.6 ± 10.6</td>
<td>18.1 ± 17.8</td>
</tr>
<tr>
<td>Rugosity (1-5 scale)</td>
<td>3.1 ± 1.0</td>
<td>3.5 ± 0.8</td>
</tr>
<tr>
<td>Width of insular shelf (m)</td>
<td>592 ± 213*</td>
<td>265 ± 25*</td>
</tr>
</tbody>
</table>

*Mean (±SD) environmental variable values presented per location and geomorphology. A two-stage nested design in ANOVA (geomorphology nested within location) specifies modal-
ities having significantly higher (a) and lower (b) values.
Halichoeres cosmetus Randall and Smith, 1982, Zebrasoma gemmata (Vaillanciennes, 1835)

The second axis, mainly related to depth, distinguishes drop offs (top of the axis) and basin blocks (bottom of the axis). Fish species that occurred mainly on basin blocks presented a higher percentage of nocturnal carnivores [16% vs. 0% on drop offs, C. guichenoti (Sauvage, 1880), Cirrhitops mascarenensis Randall and Schultz, 2008, Pararichthys arcatus (Cuvier, 1829)], while those occurring mainly on drop offs presented a greater percentage of diurnal and nocturnal planktivores [22% vs. 5% on basin blocks, Pseudanthias evansi (Smith, 1954), Chromis chrysura (Bliss, 1883), Caesio teres Seale, 1906, Nemateleotris magnifica Fowler, 1938, Naso hexacanthus (Bleeker, 1855)].

3.3. Linear regressions

Only two fish assemblage variables were significantly related to the combination of the four independent environmental variables: the percentages of browsers of sessile invertebrates and the SWIO species richness, which 38% and 35% of the variance were predicted by the models respectively (Table 2). However, if the individual contributions of predictors on the fish assemblage variables are explored, several relationships appear to be significant. Thus, the width of the insular shelf contributed significantly to the percentages of browsers of sessile invertebrates (positive relations), omnivores and diurnal planktivores and to the SWIO species richness (negative relations). In the same way, the depth was significantly related to the percentages of diurnal and nocturnal planktivores (positive relations), nocturnal carnivores and to the SWIO species richness (negative relations). Finally, the rugosity was significantly related to the percentages of piscivores (positive relation). It can be noticed that the proportion of soft substrate was not significantly related to any fish assemblage variable.

The two significant predictors of the SWIO species richness were incorporated as environmental covariates into an ANCOVA (Table 3) to assess the specific effect of geomorphologies and locations (fixed factors) on the SWIO species richness (dependent variable) regardless of the effect of depth and width of the insular shelf. The results of the model confirmed the significant effect of the two covariates on the SWIO species richness but no significant effect of the two fixed landscape factors or

### Table 2

<table>
<thead>
<tr>
<th>Fish assemblage variables</th>
<th>Depth</th>
<th>Soft substrate</th>
<th>Rugosity</th>
<th>Width of the insular shelf</th>
<th>Multiple R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herbivores</td>
<td>0.02</td>
<td>0.29</td>
<td>0.28</td>
<td>0.20</td>
<td>0.18</td>
</tr>
<tr>
<td>Omnivores</td>
<td>0.03</td>
<td>−0.29</td>
<td>−0.07</td>
<td>−0.34*</td>
<td>0.17</td>
</tr>
<tr>
<td>Browsers of sessile invert.</td>
<td>−0.13</td>
<td>0.08</td>
<td>0.34*</td>
<td>0.60**</td>
<td>0.38**</td>
</tr>
<tr>
<td>Diurnal carnivores</td>
<td>−0.28</td>
<td>0.26</td>
<td>−0.16</td>
<td>−0.12</td>
<td>0.22</td>
</tr>
<tr>
<td>Nocturnal carnivores</td>
<td>−0.37*</td>
<td>0.05</td>
<td>0.23</td>
<td>0.04</td>
<td>0.17</td>
</tr>
<tr>
<td>Piscivores</td>
<td>0.13</td>
<td>0.32</td>
<td>0.34*</td>
<td>−0.21</td>
<td>0.23</td>
</tr>
<tr>
<td>Diurnal planktivores</td>
<td>0.36*</td>
<td>0.05</td>
<td>−0.16</td>
<td>−0.37*</td>
<td>0.23</td>
</tr>
<tr>
<td>Nocturnal planktivores</td>
<td>0.41*</td>
<td>−0.07</td>
<td>0.23</td>
<td>0.04</td>
<td>0.17</td>
</tr>
<tr>
<td>SWIO species richness</td>
<td>−0.42*</td>
<td>−0.02</td>
<td>−0.13</td>
<td>−0.42*</td>
<td>0.35**</td>
</tr>
<tr>
<td>Non-SWIO species richness</td>
<td>−0.10</td>
<td>0.13</td>
<td>0.32</td>
<td>0.19</td>
<td>0.11</td>
</tr>
<tr>
<td>Total species richness</td>
<td>−0.19</td>
<td>0.13</td>
<td>0.26</td>
<td>0.07</td>
<td>0.09</td>
</tr>
</tbody>
</table>
interaction among variables. The projection of the linear regression plan of the SWIO species richness as a function of depth and width of the insular shelf on a three-dimensional plot also showed similar slopes in all landscape features (Fig. 3). In this context, it could be argued that the contribution of the landscape features to the SWIO species richness is determined by the depth and width of the island shelf, but also by the age of the lava flow and the distance to the most recent lava flow that were highly correlated with the width of the insular shelf. Therefore, stations characterized by shallow depths, narrow insular shelf, short distance to the most recent lava flow and a relatively young age are colonized by the highest SWIO species richness, whatever their geomorphology or location.

4. Discussion

4.1. Influence of selected environmental variables on fish community structure

Width of the insular shelf was the environmental variable with the most influence on the structure of fish communities (Table 2). It was strongly correlated with age of the lava flow and distance to the most recent lava flow. This combination of variables had different but related effects on fish assemblages. According to Emslie et al. (2010), the enlargement of the insular shelf may promote more abundant benthic resources and provide more shallow ecological niches. The concept of the niche predicts that more species can coexist in areas with a greater variety of resources, as there would be more potential niches to be occupied by habitat specialists (Hutchinson, 1957). These observations may explain the higher proportions of browsers of sessile invertebrates, feeding on benthic resources, on stations with a larger insular shelf and the greater percentages of diurnal planktivores and omnivores, drawing opportunistically their food from the water column, on stations with narrow insular shelf. Moreover, diurnal planktivores densely colonize environments exposed to tidal currents, along slopes adjacent to deeper water, their main prey (copepods, pelagic tunicates, fish eggs) being more accessible (Friedlander and Parrish, 1998; Hobson, 1991; Hobson and Chess, 1978). It is therefore not surprising to observe them in higher proportions on stations with a narrower insular shelf, characterized by steep slopes within the surveyed area.

Depth was also a very influential variable on the fish community structure. This influence is mainly due to changes in environmental variables such as hydrodynamism, water desalination, light intensity, oxygenation, and exogenous physicochemical inputs and its effects on the availability of food resources, some being more abundant near the surface (algae, corals, phytoplankton) and others at depth (macrobenthic invertebrates, zooplankton). These trends have been studied in many regions and contexts, including Indo-Pacific reefs (Booth and Wellington, 1998; Friedlander and Parrish, 1998), Caribbean reefs (Greenfield and Johnson, 1990; Lara and González, 1998) or rocky Mediterranean shores (Bell, 1983; Dufour et al., 1995). In this study, nocturnal and diurnal planktivores were recorded at greater depths. In addition to the ecological preferences of diurnal planktivores already mentioned, nocturnal planktivores aggregate preferentially in environments protected from wave and current. They are strict carnivores and feed on large organisms (large calanoids, mysids, isopods, amphipods, decapod larvae) (Hobson and Chess, 1978). These environmental conditions are found at greater depths, due to stringent hydrodynamism that was generated by trade wind swells on surface (Letourneur, 1998). Conversely, nocturnal carnivores were recorded in higher proportions in shallow waters. This minority diet (10% of recorded species) was mainly represented by Cirrhitidae species (51% of nocturnal carnivores occurrences, e.g. C. guichenoti, C. mascarenensis, P. arcatus), mainly recorded at depths of 20 m and less (Lieske and Myers, 1994).

It was recognized that rugose substrata may offer more shelter from predators or nesting sites. Gratwicke and Speight (2005) showed that at least 35% of the fish species observed in a range of shallow tropical marine habitats were positively related to rugosity. Moreover, the high rugosity often offered by coral cover can explain the relationship between rugose substrates and some ecological characteristics such as high percentages in corallivore organisms and high Shannon–evenness indexes (Purkis et al., 2008). These previous results can explain the relationship between rugosity and percentages of browsers of sessile invertebrates, which feed especially on coral polyps but also on other fixed organisms, observed during this study. In a previous work on underwater lava flows of the Piton de la Fournaise, Pinault et al. (2013b) also noted the positive influence of scoriaceous rocks and unconsolidated rubble and boulders of the most recent lava flow on substrate rugosity. This second origin of substrate rugosity could explain the positive relationship between this variable and the percentage of piscivores. Indeed, Godwin and Kosaki (1989) observed on the latest lava flow of the Kilauea volcano, in Hawaii, abundances of juvenile fishes much higher than on other flows. Pinault et al. (in press) also described on the lava flows of the Piton de la Fournaise a mass-settlement event of juvenile fish (e.g. Parupeneus cyclostomus (Lacepède, 1801), Pterois antennata (Bloch, 1787), Pterois

Table 3

<table>
<thead>
<tr>
<th>Source</th>
<th>df</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>p</th>
</tr>
</thead>
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<td>40.94</td>
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<tr>
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<td>0.42</td>
<td>0.08</td>
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<td>0.55</td>
<td>0.10</td>
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</tr>
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<td>0.59</td>
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<tr>
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<td>2.53</td>
<td>0.45</td>
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<td>0.14</td>
<td>0.03</td>
<td>0.87</td>
</tr>
<tr>
<td>Depth × location</td>
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<td>1.39</td>
<td>0.25</td>
<td>0.62</td>
</tr>
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<td>0.19</td>
<td>0.19</td>
<td>0.03</td>
<td>0.86</td>
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<td>0.96</td>
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<td>5.59</td>
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<td>128.13</td>
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</table>

Figure 3. Three dimension plot of the southwestern Indian Ocean species richness as a function of width of the insular shelf and depth. □, basalt blocks; ○, drop-offs; ●, dark dots, outside volcanic enclosure; clear dots, inside volcanic enclosure. Coefficient of determination of the overall straight-line regression ($R^2$) on width of the insular shelf = 0.14 and on depth = 0.24.
synchronous species, i.e. 81%, were at maximum depth of 40 m), with high fecundity (23 species, i.e. 74%, had a minimum population doubling time of 30 cm), encountered in shallow waters (25 species, i.e. 81%, were at maximum depth of <40 m), with high fecundity (23 species, i.e. 74%, had a minimum population doubling time of <15 months), non-selective diet (10 species, i.e. 32%, were omnivores), and narrow species range-size (22 species, i.e. 71%, were sedentary). The relatively high SWIO species richness observed on the shallower stations with narrower insular shelf, younger age and located closer to the most recent lava flow could be thus the result of these convergent ecological traits, which are characteristic of species commonly found in convergent ecotopes of the impacted communities. The relative distance from the most recent lava flow in Hawaii to be about 20 to 50 years, depending on the succession stage of the impacted communities. The relative distance from the most recent lava flow of the areas outside VE could thus allow the conservation of structured reef communities, as areas subject to direct volcanic impacts would be characterized by constantly interrupted successions, resulting in pioneer stages.

Drop-offs outside VE are also relatively protected from the influence of the most recent lava flows. However, their location in deeper water attenuates the influence of swell, tidal currents and light energy on biological communities. These conditions limit the development of coral communities, but favor the aggregation of nocturnal planktivores, adapted to low hydrodynamic environments (Hobson and Chess, 1978).

Basalt blocks inside VE are distinguished by their proximity to the most recent lava flows and their regular acute volcanic disturbances. Their shallowness also exposes them to stringent environmental variables such as swell, tidal currents or chronic freshwater inputs. Moreover, the narrowness of the insular shelf, characterized by steep slopes, reduces number of shallow ecological niches and leads to regular food shortages, particularly from benthic resources (Emeslie et al., 2010). This very rigorous environment is conducive to colonization by rather broad food spectrum species like omnivores, but also to small nocturnal species.
carnivores of the Cirrhitidae family, which find shelter in the numerous topographic irregularities created by scoriaceous rocks and unconsolidated blocks from the most recent lava flows. These conditions are also suitable for colonization by the SWIO species, adapted to changing environments and frequent food shortages, compensating their low competitiveness by their high demographic and environmental flexibility (Odum, 1969; Ramade, 2003).

Finally, drop-offs inside VE are differentiated from basalt blocks by their greater depths, removing them from the direct influence (Odum, 1969; Ramade, 2003). Thus, favoring the colonization of these habitats by planktivores.

4.4. Conclusions

This study constitutes one of the few works conducted on the colonization of recent lava flows by rocky shore fish communities in coral reef environment. Overall, it addresses the mechanisms of natural selection and speciation of endemics from ecological and behavioral analyzes and not from genetic tools, very frequently used when it comes to evolutionary biology and cryptic species (e.g. Cowman and Bellwood, 2013; DiBattista et al., 2013; Hubert et al., 2012). This empirical approach provides avenues for original interpretation; particularly concerning environmental forces and constraints involved in the selection mechanisms in the SWIO region. It appears that acute pressures of volcanic origin and their direct impact on the abundance and availability of food resources could generate harsh environmental conditions, creating ecological niches weakly coveted by biological communities. This low competition for habitat could then promote the emergence of opportunistic species, especially characterized by a strong demographic and dietary flexibility (Odum, 1969; Ramade, 2003).

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