Influence of Freshwater Discharges on Biogeochemistry and Benthic Communities of a Coral Reef Ecosystem (La Réunion Island, Indian Ocean)

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We investigated the influence of freshwater discharges on biogeochemistry and benthic communities of La Saline fringing reef, the largest coral reef system of La Réunion Island (France, Indian Ocean). Sampling was conducted in February–March 2012 along a land-ocean continuum, including rivers, groundwater, reef waters, and oceanic waters. Our results indicated the discharges of basalt groundwater (enriched in nitrates) in the southern part of the reef, and the discharges of sand groundwater and river waters (enriched in organic matter) in sites located northward. These discharges mainly impacted fore reef surface waters, while groundwater resurgences were more diffuse and not clearly detected in depth. We highlighted the relationship Si(OH)4/NO3− to identify basalt groundwater discharges. We also put forward the relationships humic-like fluorescence/NH4+ and tryptophan-like fluorescence/PO43− to identify sand groundwater discharges and wastewater pollutions, respectively. We observed the eutrophication of the southern part of the reef through higher phytoplankton biomasses, a massive presence of fleshy algal formations and a low coral coverage. The spatial homogeneity of the benthic community structures onto the entire slope contrasted with the spatial heterogeneity of chemical variables in water. This reflected the different spatial and temporal scales involved in biological and chemical processes in coral reef ecosystems.

Keywords: coral reefs, La Réunion, Indian Ocean, eutrophication, groundwater, nutrients, organic matter, fluorescence

INTRODUCTION

Coral reefs are ecosystems of great biological productivity and diversity (Fisher et al., 2015) that are present mostly in shallow coastal waters of oligotrophic tropical and subtropical oceans (Kleypas et al., 1999; Hoegh-Guldberg et al., 2017). They are recognized as being particularly vulnerable to anthropogenic pressures that are exerted both through the direct exploitation of reef resources and
the multiple human activities on land, which lead to impacts at the global and local levels (Dubinsky and Stambler, 1996; Pandolfi et al., 2011; Ban et al., 2014; Hughes et al., 2017). These disturbances have caused an extensive degradation of coral reefs, which, in many cases, have undergone a phase shift from coral to algal dominance (Hughes, 1994; McManus and Polungen, 2004; Norström et al., 2009; Bonaldo and Bellwood, 2011).

Land-based inputs of freshwater are substantial sources of nutrients, organic matter (OM) and pollutants to coastal reefs (Fabricius, 2005; Tanaka et al., 2013; Lubarsky et al., 2018). Among these freshwater inputs, submarine groundwater discharge (SGD) has an important contribution, especially in tropical islands, where it can supply large amounts of terrestrial-derived nutrients to reef waters (Lewis, 1987; Paytan et al., 2006; Kim et al., 2011; Moosdorf et al., 2015). Also, in some volcanic islands of the tropical Pacific and Indian Oceans, SGD may exceed river inputs and thus represent the largest—or the only—source of nutrients to coral reefs (Street et al., 2008; Cuet et al., 2011; Lee and Kim, 2015; Lubarsky et al., 2018). Groundwater is generally enriched in dissolved nitrogen (mainly nitrates) relative to phosphorus, displaying high N/P ratios (Slomp and Van Cappellen, 2004; Street et al., 2008; Santos et al., 2013; McMahon and Santos, 2017). Because reef waters are often oligotrophic and N-limited, SGD may lead to their eutrophication, which is characterized by the shift from corals to algal communities (Koop et al., 2001; Dailer et al., 2010; Zubia et al., 2018).

Besides nutrients and other inorganic species, SGD and rivers may also carry dissolved organic matter (DOM) to coral reefs. Recent works have shown significant inputs of groundwater-derived chromophoric and fluorescent DOM (CDOM and FDOM, respectively) in various coastal reef environments (Tedetti et al., 2011; Nelson et al., 2015; Kim and Kim, 2017; Webb et al., 2019). Humic-like fluorophores have been identified as major components of groundwater FDOM, and it has been proposed to use FDOM to differentiate groundwater types and to track their dispersal into reef waters (Tedetti et al., 2011; Nelson et al., 2015; Kim and Kim, 2017). In addition to direct allochthonous inputs, an autochthonous production of FDOM may occur in the reef waters due to the SGD-induced eutrophication or other processes (Blanco et al., 2008; Tedetti et al., 2011; Nelson et al., 2015). CDOM/FDOM in coral reef waters could influence the reef biota by supplying carbon, but also by modifying the seawater optical properties and subsequently the attenuation (and availability) of solar UV and visible radiations in the water column (Coble, 2007; Nelson and Siegel, 2013; Sempère et al., 2015; Martías et al., 2018).

Even though numerous coral reefs of volcanic islands are strongly impacted by SGD, the effects of the latter on the reef water chemistry at a large spatial scale remain little known so far, particularly concerning DOM descriptors. Moreover, the potential link between the distribution of biogeochemical/chemical parameters of reef water and the distribution of benthic community structures has rarely been assessed (Blanco et al., 2008; Li et al., 2015; Naumann et al., 2015).

La Saline fringing reef is the most important coral reef complex of La Réunion Island (France, southwestern Indian Ocean). SGD from volcanic aquifer has been shown to be the main source of freshwater to La Saline reef (Jion et al., 1988) and to enrich its waters with nitrates derived from anthropogenic sources (Cuet et al., 1988; Naîm, 1993; Mioche and Cuet, 1999). Several studies have been conducted so far on this ecosystem to determine the effects of SGD/eutrophication on C, N, P and O fluxes (Clavier et al., 2008; Cuet et al., 2011; L’Helguen et al., 2014), the distribution of FDOM and hydrocarbons (Tedetti et al., 2011; Guigue et al., 2015), the rates of bioerosion (grazers/borers) and calcification (corals) (Chazottes et al., 2002; Chauvin et al., 2011), and the distribution of coral and algal communities (Tourrand et al., 2013; Zubia et al., 2018). Nevertheless, since most of these works have focused on quite limited geographical (back reef) areas of the reef, the distribution of chemical and biological parameters at a larger spatial scale (from back to fore reef waters, from north to south of the reef) has not been evaluated yet.

In this context, the objectives of the present work are (1) to assess the distribution of both nutrients and DOM/FDOM descriptors at the whole reef scale, i.e., from north to south, and along a continuum from land to ocean, including rivers, groundwater discharges, reef waters and oceanic waters. (2) To highlight relevant indicators of freshwater discharges into back reef and fore reef waters. (3) To attempt linking the distribution of chemical parameters with phytoplankton biomass and benthic community structure.

**MATERIALS AND METHODS**

**Study Area**

Extending over 9 km long, up to 500 m wide, and from 1 to 1.5 m deep, La Saline fringing reef is the largest coral complex of La Réunion Island (Cordier, 2007; L’Helguen et al., 2014). From land toward oceanic waters, it includes two principal water complexes: (1) the back reef area, located within the back reef zone and covered with sandy sediment (mean depth: 1.2 m) and the internal reef flat area characterized by transverse strips colonized by branched corals (mean depth: 0.5 m), and (2) the fore reef waters, located close to the outer reef slope dominated by corals and macroalgae. The fore reef compartment gently slopes from the reef flat environment, subjected to strong swells and waves, down to 20-m deep (Figure 1).

Since the 1980s, it has been observed a degradation of La Saline ecosystem through a constant decrease in the living coral cover correlated with a raise in algal cover in the reef flat zone and on the outer reef slope (Ahamada et al., 2008; Naîm et al., 2013; Zubia et al., 2018; Nicet et al., 2019). Although varying spatially and seasonally (Cuet et al., 2011; Naîm et al., 2013), algal cover overall has displayed a substantial increase since the late 1990s in La Saline fringing reef (Tourrand et al., 2013). This eutrophication has been in part attributed to a chronic enrichment in nitrates derived from SGD (Cuet et al., 1988; Naîm, 1993; Naîm et al., 2013).

Groundwater regular discharges are noticeable along most of La Saline shoreline (Cuet et al., 1988; Naîm et al., 2000). These discharges vary from north to south of the reef depending on the thickness of the coral detrital-based sedimentary formations.
FIGURE 1 | (A) Location of the study stations in La Saline fringing reef system (La Réunion Island, southwestern Indian Ocean), including rivers, groundwater (freshwaters), back reef waters, fore reef waters, and oceanic waters. “North” includes fore reef sites located north of the Hermitage river (i.e., SGF, PMF, TF, HF), while “South” gathers fore reef sites positioned south of the Hermitage river (i.e., CFF, PAF, BSF, SSF, 3BF). Station map made from Google Earth Pro V 7.3.0.3832 (August 18, 2017). The detailed characteristics of these stations are provided in Supplementary Table S1. (B) Cross (conceptual) section of one transect (example for Planch’Alizés transect, PA) showing the different water bodies sampled from back reef to oceanic waters, and the potential discharges of sand groundwater into back reef waters, basalt groundwater into back reef and fore reef waters, as well as the water transfer from back reef to fore reef.

(sand layer groundwater, SL), which lie over the highly permeable volcanic aquifer (basalt layer groundwater, BL) (Join et al., 1988; Leze, 2012). In the central part of La Saline (Hermitage area), sand layer is the thickest, preventing the discharge of basalt groundwater in the back reef area, but allowing possibly its release beyond the reef front, i.e., in the fore reef waters. The thickness of the sand layer gradually decreases southward. Hence, in the southernmost part of La Saline (BS4-Trois-Bassins area) where sand layer has almost disappeared, basalt groundwater outcrops directly flows in the back reef area. In addition to groundwater, three rivers are freshwater sources for the reef: The Hermitage River (HR) (in the central part), receiving a treated wastewater effluent, the Saint Gilles River (SGR) (in the north part), whose mouth hosts a marina, and the Trois-Bassins River (3BR; in the south part) (Figure 1 and Supplementary Table S1). Besides, around 60% of habitations in this area have septic tank sewage systems. This water discharges directly into the unsaturated zone which then percolates into groundwater (Rogers et al., 2012).
Hydrodynamics in La Saline fringing reef is constrained by the reef geomorphology, the open ocean forcing (tide and waves), and wind (Cordier et al., 2013). Water circulation in the reef is structured in several individual cells (Cordier, 2007). In each of those cells, wave breaking near the reef crest drives a water flow across the reef flat up to the back reef area where water progresses along the shoreline either toward the north end of the reef (Saint Gilles area), the Hermitage pass (in the central part of the reef, north and south of the Hermitage River) or toward the Trois-Bassins pass (in the south) (Figure 1). The reef water flows back to open sea mainly through these two passes but also via narrow gaps in the reef crest. Water residence time in the back reef zone varies from ~2 h 30 min to 10 h 30 min, and tides are semidiurnial ranging from 0.1 m (neap) to 0.9 m (spring) (Cordier, 2007).

**Sampling and in situ Measurements**

Sampling was conducted in freshwaters, back reef waters, fore reef waters and oceanic waters from February 22, to March 13, 2012 in the morning during the summer season, which is generally characterized by high seawater temperatures and strong rains (Figure 1 and Supplementary Table S1). Water sampling for measurements of salinity and biogeochemical parameters (nutrients, DOM and pigments) was carried out in all these water bodies, while sampling of benthic communities was performed only onto outer reef slopes (fore reef waters). All water samples were retrieved in 4-L Nalgene® polycarbonate bottles extensively washed with 1 M hydrochloric acid (HCl) and ultrapure water (i.e., Milli-Q water from Millipore system, final resistivity of 18.2 MΩ cm) before use. The bottles were rinsed three times with the respective sample before filling and placed in the cold and in the dark after collection until back to the laboratory. Also, in situ measurements of temperature, salinity and total chlorophyll a (TChl a) were undertaken in most of back reef, fore reef, and oceanic sites (Supplementary Table S1) using a 19plus conductivity-temperature-depth (CTD) probe (Sea-Bird Electronics, Inc.) equipped with a WETStar fluorometer (WETLabs, Inc.). The detailed sampling for the different water bodies is provided in Supplementary Text S1.

**Filtration of Water Samples and Storage**

Back in the laboratory, water samples were immediately filtered under a low vacuum. Samples for dissolved organic carbon (DOC) and FDOM analyses were filtered through pre-combusted (500°C, 4 h) glass fiber filters (GF/F) (47-mm diameter, Whatman). The ~0.7-µm filtered samples were stored at ~18°C until analysis. Samples for the determination of salinity and nitrate (NO$_3^-$), nitrite (NO$_2^-$), ammonium (NH$_4^+$), phosphate (PO$_4^{3-}$), dissolved organic nitrogen (DON), and urea concentrations were filtered through pre-combusted GF/F filters (47-mm diameter, Whatman), and those for the determination of silicate [Si(OH)$_4$]$_a$ concentration, through 0.45-µm mixed cellulose ester filters (25-mm diameter, HAWP Millipore). Filtered samples were stored in 125-mL Nalgene® polycarbonate bottles at ~18°C until analyses, except those for NH$_4^+$ (stored in 100-mL Schott® glass bottles), Si(OH)$_4$ (stored at 4°C) and salinity (stored in sealed cans in the dark at room temperature). Samples for the determination of TChl a and phaeopigment a (Pheo a) concentrations were filtered (from 50 to 400 mL for freshwater samples to 1000 mL for marine samples) through pre-combusted GF/F filters (25-mm diameter, Whatman). Filters were stored at ~18°C until analyses.

In this work, TChl a and Pheo a concentrations were not determined in groundwater (i.e., SL1, 2, BL1-3, TE, FH, FM samples), while, NH$_4^+$, PO$_4^{3-}$, DON, urea, TChl a, and Pheo a concentrations were not determined in back reef waters, except in HB (Supplementary Table S1). For back reef waters, TChl a concentrations derived from CTD measurements were used.

**Measurements of Salinity and Biogeochemical Parameters**

**Salinity**

Salinity was determined on all discrete samples in the laboratory using a calibrated Yeo-Kal Mark IV salinometer. The accuracy of the salinity analysis was estimated ±0.003.

**Nutrients, DON and Urea**

Concentrations in nutrients, DON and urea were determined by colorimetric methods. Si(OH)$_4$, NO$_3^-$, NO$_2^-$, and PO$_4^{3-}$ were analyzed on an automatic analysis chain (Autoanalyzer III Seal) according to standard methods by Tréguer and LeCorre (1975) and modified by Aminot and Keroulé (2007) for PO$_4^{3-}$. Precision of measurements was ±0.05 µM for Si(OH)$_4$ and NO$_3^-$, and ±0.02 µM for NO$_2^-$ and PO$_4^{3-}$. NH$_4^+$ and urea were analyzed with a UVIKON 922 spectrophotometer equipped with 10-cm cuvettes according to methods by Aminot (1983) for NH$_4^+$, and Mulvenna and Savidge (1992) and Goeyens et al. (1998) for urea. Precision of measurements was ±0.05 and ±0.03 µM for NH$_4^+$ and urea, respectively. Total dissolved N was determined by the wet oxidation procedure described in Raimbault et al. (1999), and DON concentration was retrieved by subtracting NO$_3^-$ + NO$_2^-$ + NH$_4^+$ concentration to total dissolved N concentration. Precision of measurements was ±5%.

**TChl a and Pheo a**

Total chlorophyll a and Pheo a concentrations were determined by fluorescence after methanol extraction according to Raimbault et al. (2004), by using a Turner Fluorometer 110 equipped with the Welschmeyer kit to avoid chlorophyll b interference (Welschmeyer, 1994). The detection limit of TChl a was ~0.01 µg L$^{-1}$. Calibrations were made using a pure Sigma Chl a standard. Pheo a concentration was obtained by difference between TChl a concentrations measured before and after acidification.

**DOC**

For DOC analysis, the samples were spiked with phosphoric acid and bubbled for 10 min with CO$_2$-free air to purge inorganic carbon. DOC was measured on two replicates by high-temperature catalytic oxidation using a Shimadzu TOC 5000 Total Carbon Analyzer (Sohrin and Sempéré, 2005). The accuracy and system blank of the instrument were determined by the analysis of Deep Atlantic Water and low carbon water reference standards. The nominal precision of the analysis procedure was within 2%.
FDOM
Fluorescent DOM analyses were carried out with a Hitachi F-7000 Fluorescence Spectrophotometer. The method is detailed in Tedetti et al. (2012, 2016) and Ferretto et al. (2017). In brief, excitation-emission matrices (EEMs) were produced for excitation wavelengths (\(\lambda_{\text{Ex}}\)) between 200 and 500 nm and for emission wavelengths (\(\lambda_{\text{Em}}\)) between 280 and 550 nm. Two replicates were run for each sample. Inner filtering effects were corrected by applying the absorption method (Ohno, 2002). Absorbance measurements were performed between 200 to 800 nm with a Shimadzu UV-Visible 2450 Spectrophotometer.

The EEMs were then blank-corrected by subtracting the pure water EEM, and converted into quinine sulphate unit (QSU). EEMs were then processed using parallel factor analysis (PARAFAC), operated using the DOMfluor toolbox v1.6, running under MATLAB 7.10.0 (R2010a). The datasets of freshwaters (rivers, groundwater) and marine waters (back reef, fore reef, and oceanic waters) were processed separately due to their differences in salinity values and fluorescence intensities (Stedmon and Bro, 2008; Ferretto et al., 2017). Thirty samples (EEMs) were used for the freshwater PARAFAC model, while 91 samples were used for the marine water PARAFAC model.

In this study, we validated two PARAFAC components for each modeled dataset (freshwater and marine samples): one humic-like component (\(\lambda_{\text{Ex}1}, \lambda_{\text{Ex}2}/\lambda_{\text{Em}}\) of 230, 315/440 nm for freshwater samples, and \(\lambda_{\text{Ex}1}, \lambda_{\text{Ex}2}/\lambda_{\text{Em}}\) of 235, 315/436 nm for marine samples) and one tryptophan-like component (\(\lambda_{\text{Ex}1}, \lambda_{\text{Ex}2}/\lambda_{\text{Em}}\) of 230, 290/348 nm for freshwater samples, and \(\lambda_{\text{Ex}1}, \lambda_{\text{Ex}2}/\lambda_{\text{Em}}\) of 225, 280/340 nm for marine samples). These two components (humic- and tryptophan-like) are common and ubiquitous FDOM fluorophores classically found in aquatic environment (Coble, 2007; Fellman et al., 2010).

Benthic Community Sampling and Analysis
For each site, samples for the analysis of benthic communities were taken by divers onto the outer reef slope at 10-m depth (samples n° 4 for all fore reef sites) (Supplementary Table S1). Line Intercept Transect (LIT) developed by English et al. (1994) was adapted for the monitoring of coral reef in the Indian Ocean (Conand et al., 1997, 2000). LIT was used to characterize the benthic community structure at an expert level for the coral species (hard and soft corals), the main algal assemblages (i.e., coralline algae, macroalgae, turf algae) and the abiotic substrates. Three replicates of 20-m long were recorded for each station.

Statistical Treatments
In most cases non-parametric tests were preferred to parametric tests (Joliffe, 1986). Principal component analyses (PCA) were performed based on Spearman’s rank order correlation matrices. In some cases, Pearson correlations were applied when the slope and y-intercept of the linear regression were investigated. Hierarchical ascendant classification (HAC) was used to find homogenous groups of samples within the dataset, with respect to selected variables. The dissimilarity measurement between groups was based on Ward's method, which uses squared Euclidean distance. Variable values were centered and reduced beforehand. Non-parametric tests of Mann–Whitney (U-test) and Kruskal–Wallis (H-test) were used to compare the distributions of two (U-test) or more than two (H-test) groups. Normality tests, PCA, HAC, correlations, box-and-whisker plots, U- and H-tests were carried out on salinity and all biogeochemical parameters (nutrients, DOC, DON, urea, FDOM, TChl a, and Pheo a) using XLSTAT 2013.5.01. Moreover, canonical correspondence analysis (CCA) was performed with PAST 3.25 software, following the eigen analysis algorithm given in Legendre and Legendre (1998) to investigate the potential influence of biogeochemical parameters on the distribution of benthic communities. CCA was applied on salinity, nutrients, FDOM, and benthic communities (abundances of mixed algal assemblages, coralline algae, Acropora corals, non-Acropora corals, soft corals, abiotic substratum).

Color maps of CTD data were carried out using Ocean Data View (ODV) software version 4.6.5 (Schlitzer, 2014). The spatial interpolation/gridding of data was conducted using Data-Interpolating Variational Analysis (DIVA) (Barth et al., 2010; Troupin et al., 2012).

RESULTS

Freshwaters
Salinity and Biogeochemical Parameters
Salinity and biogeochemical parameters in freshwater samples (rivers and groundwater) are presented Figure 2 and Supplementary Table S2. Basalt layer groundwater (BL) was enriched in Si(OH)\(_4\) (850–1250 µM) and NO\(_3^-\) (270–800 µM) compared to other freshwater bodies (Figures 2B,C), while displaying very low contents in reduced forms of dissolved inorganic nitrogen (0.03–0.08 µM for NO\(_2^-\) and 0.00–0.17 µM for NH\(_4^+\)) and DOM, including DOC (25–45 µM), DON (0.0–7.4 µM), urea (0.04–0.49 µM), humic- and tryptophan-like fluorophores (5–80 QSU) (Figures 2D,E,G–K).

Samples from Trois-Bassins River (3BR) were brackish with salinity ranging from 26.0 to 27.9, and exhibited low concentrations in Si(OH)\(_4\) (125–150 µM) and PO\(_4^{3-}\) (0.19–0.22 µM) relative to other freshwater bodies, as well as low concentrations in NO\(_3^-\) (5.6–15.7 µM) (Figures 2A–C,F). On the other hand, 3BR was enriched in DOM (DOC: 270–340 µM; humic-like: 660–1040 QSU), and displayed higher concentrations in DON (10.2–14.4 µM), NO\(_2^-\) (∼1 µM), and NH\(_4^+\) (22.8–31.5 µM) compared to BL (Figures 2D,E,G,H,J).

Sand layer groundwater (SL) presented salinities of 5.7–6.0, and contents in DOM, NO\(_3^-\), and NH\(_4^+\) of the same order of magnitude that those of 3BR. Also, concentrations in Si(OH)\(_4\) (580–620 µM) and PO\(_4^{3-}\) (∼1.2 µM) were lower than in BL, while NO\(_3^-\) had almost disappeared (0.08–1.51 µM) (Figures 2A–K).

Hermitage River (HR) had high contents in DOC (∼390 µM) and humic-like material (890–1095 QSU) of the same order of magnitude than those of SL and 3BR. However, compared to these two water bodies, it showed a significant enrichment in
DON (\(\sim 42 \mu M\)), tryptophan-like fluorophore (\(\sim 935\) QSU), and in a lesser extent, urea (\(\sim 0.85 \mu M\)), PO\(_4^{3-}\) (\(\sim 3.6 \mu M\)), and NO\(_2^-\) (\(\sim 2.2 \mu M\)). In contrast, NH\(_4^+\) concentration in HR (2.4–5.8 \(\mu M\)) was much lower than in SL and 3BR (Figures 2D–K).

Saint Gilles River was distinguished from other rivers and SL due its lower content in DOM, particularly in DOC (\(\sim 60 \mu M\)) and humic-like material (\(\sim 100\) QSU) (Figures 2G–K).

**PCA Discrimination of Samples**

Spearman PCA applied on salinity and biogeochemical parameters for freshwater samples are shown Supplementary Figure S1. Correlation circle displayed strong positive (linear) correlations between Si(OH)\(_4\), NO\(_3^-\), and PO\(_4^{3-}\) (\(r = 0.78–0.91\)), between NO\(_2^-\), DOC, DON, humic-, and tryptophan-like fluorophores (\(r = 0.80–0.94\)), between humic-like and NH\(_4^+\) (\(r = 0.80\)), and between DOC and urea (\(r = 0.78\)) (\(n = 15, p < 0.05\) for all \(r\) values).

The first principal component (PC1), which explained 72.9% of total variability within samples, allowed the discrimination between (1) BL and SGR, which were (especially BL) enriched in Si(OH)\(_4\), NO\(_3^-\), and PO\(_4^{3-}\) but depleted in DOM and reduced forms of dissolved inorganic nitrogen, and (2) SL, 3BR, HR, which presented higher contents in DOM and NO\(_2^-\)/NH\(_4^+\) than BL, as well as lower concentrations of Si(OH)\(_4\) and NO\(_3^-\). The second principal component (PC2) accounted for 12.5% of total variability, and allowed separating HR from SL and 3BR due to
the enrichment of HR in DON, tryptophan-like material, urea, PO$_4^{3-}$ and NO$_3^-$ . The three groups of samples highlighted by PCA (G1: SGR and BL, G2: BR and SL, G3: HR) were confirmed by the HAC (Supplementary Figure S1).

**Back Reef Waters**

**Distribution of Salinity and Biogeochemical Parameters**

The distribution of salinity and biogeochemical parameters in back reef waters are shown Figure 3, Supplementary Table S2, and Supplementary Figure S2. For all sites, except the northmost one (TB), we observe gradients from the beach to the reef flat, with increasing salinity values (for instance from 33.49 to 35.62 for TEB) and decreasing Si(OH)$_4$, NO$_3^-$, and NO$_2^-$ concentrations (for example from 65 to 2 µM of Si(OH)$_4$ for TEB). North-South differences also appeared with overall lower salinities and higher nutrient concentrations in the southern sites (PAB, TEB, GTB), especially in TEB, relative to the northern ones (TB, CMB). These North–South differences were particularly marked when considering the beach stations (Figures 3A–D). The distributions of Si(OH)$_4$ and NO$_3^-$ across sites were quasi superimposed. DOC concentrations, which did not present any beach-reef flat gradient, were maximal at PAB site (86–99 µM) (Figure 3E). The distribution of humic-like fluorophore was close to that of NO$_2^-$ . Indeed, the humic-like content was minimal at TB (5.5–6.5 QSU) and maximal at GTB-beach (24 QSU), while displaying a progressive decrease from the beach to the reef flat at TEB (Figure 3F). Tryptophan-like material showed the highest values and variability in TEB and GTB sites (22–147 QSU) (Figures 3A–D). The distribution of salinity and biogeochemical parameters in fore reef and oceanic waters is presented in Supplementary Figures S5–S7. In addition, the detailed description for each parameter is provided in Supplementary Text S2. Salinity, PO$_4^{3-}$-concentration and fluorescence intensity of tryptophan-like fluorophore were significantly higher in the northern part (from SFG to HF) than in the southern part of the reef (from CFF to 3BF) (U-test, p < 0.05) (Figures 4A,F,K and Supplementary Figures S5D–F, S6C,R, S7O). On the contrary, Si(OH)$_4$, DOC and TChl $a$ concentrations were significantly higher in the south than in the north (U-test, p < 0.05) (Figures 4B,G,L and Supplementary Figures S5G–I, S6F, S7C,R). All the other parameters did not show any difference between the north and the south (U-test, p > 0.05). Salinity displayed significantly lower values in surface waters than in waters close to the slope (U-test, p < 0.05), while the inverse pattern (significantly higher values in surface waters) was found for Si(OH)$_4$, NO$_3^-$, NO$_2^-$ and NH$_4^+$ concentrations, as well as fluorescence intensity of humic-like fluorophore (U-test, p < 0.05) (Figures 4A–E, and Supplementary Figures S5E,J, S6H,K,N, S7K). The other parameters did not present any difference between surface and slope (U-test, p > 0.05). For salinity and Si(OH)$_4$ concentration, no particular gradient was observed as a function of the distance to the reef front (i.e., between samples collected over depths of 5, 10, or 20 m) (U-test, p > 0.05) (Supplementary Figures S6A,D), whereas NO$_3^-$, NO$_2^-$, NH$_4^+$, PO$_4^{3-}$, and TChl $a$ concentrations significantly decreased with distance to the reef front (U-test, p < 0.05) (Figures 4C–F,L and Supplementary Figures S5G–I, S6G,J,M,P, S7P). All the other parameters did not significantly vary with respect to the distance to the reef front, except DON, whose concentration increased from the front to the open sea (Figure 4H and

**Fore Reef and Oceanic Waters**

**Distribution of Salinity and Biogeochemical Parameters**

The distribution of salinity and biogeochemical parameters in fore reef and oceanic waters is presented Figure 4, Supplementary Table S2, and Supplementary Figures S5-S7. In addition, the detailed description for each parameter is provided in Supplementary Text S2. Salinity, PO$_4^{3-}$-concentration and fluorescence intensity of tryptophan-like fluorophore were significantly higher in the northern part (from SFG to HF) than in the southern part of the reef (from CFF to 3BF) (U-test, p < 0.05) (Figures 4A,F,K and Supplementary Figures S5D–F, S6C,R, S7O). On the contrary, Si(OH)$_4$, DOC and TChl $a$ concentrations were significantly higher in the south than in the north (U-test, p < 0.05) (Figures 4B,G,L and Supplementary Figures S5G–I, S6F, S7C,R). All the other parameters did not show any difference between the north and the south (U-test, p > 0.05). Salinity displayed significantly lower values in surface waters than in waters close to the slope (U-test, p < 0.05), while the inverse pattern (significantly higher values in surface waters) was found for Si(OH)$_4$, NO$_3^-$, NO$_2^-$ and NH$_4^+$ concentrations, as well as fluorescence intensity of humic-like fluorophore (U-test, p < 0.05) (Figures 4A–E, and Supplementary Figures S5E,J, S6H,K,N, S7K). The other parameters did not present any difference between surface and slope (U-test, p > 0.05). For salinity and Si(OH)$_4$ concentration, no particular gradient was observed as a function of the distance to the reef front (i.e., between samples collected over depths of 5, 10, or 20 m) (U-test, p > 0.05) (Supplementary Figures S6A,D), whereas NO$_3^-$, NO$_2^-$, NH$_4^+$, PO$_4^{3-}$, and TChl $a$ concentrations significantly decreased with distance to the reef front (U-test, p < 0.05) (Figures 4C–F,L and Supplementary Figures S5G–I, S6G,J,M,P, S7P). All the other parameters did not significantly vary with respect to the distance to the reef front, except DON, whose concentration increased from the front to the open sea (Figure 4H and
Supplementary Figure S7D). It should be noticed that the spatial distribution of Pheo a, not shown in Figure 4, was analogous to that of TChl a.

PCA Discrimination of Samples
Spearman PCA applied on salinity and biogeochemical parameters for the fore reef and oceanic samples are shown in Supplementary Figure S8. Correlation circle mainly revealed a significant negative correlation between salinity and Si(OH)$_4$ ($r = -0.62$), and significant positive correlations between TChl a and Pheo a ($r = 0.86$), between Si(OH)$_4$ and NO$_3^-$ ($r = 0.60$), between NO$_3^-$, NO$_2^-$, NH$_4^+$, and PO$_4^{3-}$ ($r = 0.55–0.88$), and between NH$_4^+$, humic-like and urea ($r = 0.54–0.69$) ($n = 57$, $p < 0.05$).
First principal component accounted for 36.6% of total variability within samples. It was mostly driven by NO$_3^-$, NO$_2^-$, and NH$_4^+$, that were the only three parameters that both decreased when the distance to the reef front increased, and were higher at the surface than onto the slope. Therefore, from the left to the right side of PC1 (i.e., increasing concentrations in NO$_3^-$, NO$_2^-$, NH$_4^+$), samples present a distribution from oceanic waters and bottom/distant fore reef waters toward surface/closer fore reef waters. Humic-like and urea are significantly correlated with NH$_4^+$, hence also contributed to PC1, although more weakly (Supplementary Figure S8).

Second principal component explained 20.8% of total variability. It was principally associated (negatively) with salinity and tryptophan-like fluorophore, and (positively) with TChl a and Pheo a. It allowed the discrimination of samples mainly regarding the different sites and the north-south gradient. From the bottom to the top of PC2, i.e., increasing concentrations in TChl a and Pheo a, decreasing values of salinity and tryptophan-like, samples are roughly distributed from northern (HF, PMF, SGF, TF) to southern sites (CFF, PAF, BSF, SSF, 3BF) (Supplementary Figure S8).

Pearson Relationships
We investigated the Pearson relationships between some selected parameters in each site independently to help identify the locations and the type of freshwater inputs, as well as to highlight markers of terrestrial/anthropogenic inputs (Supplementary Table S5 and Figure 5). Negative correlations between salinity and nutrients or humic-like were the highest in the southernmost sites (BSF, SSF, 3BF), which displayed particularly strong
correlations between salinity and Si(OH)$_4$, NO$_3^-$, or NO$_2^-$ ($r = -0.92$–0.97, $n = 9$, $p < 0.05$) (Supplementary Table S5 and Figure 5A). Besides the southernmost part of the reef, PMF, HF, and PAF sites also showed significant negative correlations between salinity and Si(OH)$_4$, while SGF, TF, and CFF did not exhibit any significant correlations between salinity and the other variables, except CFF for tryptophan-like (Supplementary Table S5). Likewise, the highest correlations between Si(OH)$_4$ and NO$_3^-$ concentrations were recorded for the southernmost sites (BSF, SSF, 3BF) ($r = 0.90$–0.98, $n = 9$, $p < 0.05$) (Supplementary Table S5 and Figure 5B). The linear regressions presented slope values (1.7–4.7) of the same order of magnitude than the Si(OH)$_4$/NO$_3^-$ ratio observed in BL (2.4 ± 0.7).

The correlations between NO$_3^-$, NO$_2^-$, NH$_4^+$, and humic-like fluorescence were significant for all sites ($r = 0.64$–0.99, $n = 9$, $p < 0.05$) (Supplementary Table S5), with the highest correlations found for 3BF (NO$_3^-$ vs. NO$_2^-$; Figure 5C), PAF (NO$_3^-$ vs. NH$_4^+$; Figure 5D), or HF/PAF (NH$_4^+$ vs. humic-like; Figure 5E). Concerning the relationships between NH$_4^+$ and humic-like fluorescence, the regression slope values for sites HF
and PAF (37 and 45, respectively) (Supplementary Table S5 and Figure 5E) were in the same range than the humic-like/NH$_4^+$ concentration ratio found in freshwaters containing significant amounts of humic matter, except the Hermitage River (i.e., SL, SGR, and 3BR; ratios: 16–69).

Regarding the relationships between PO$_4^{3-}$ concentration and tryptophan-like fluorophore, the highest correlations were observed for SGF, PMF, and HF sites, all located in the northern part of the reef ($r = 0.70–0.81$, $n = 9$, $p < 0.05$) (Supplementary Table S5 and Figure 5F).

**Benthic Community Structure**

The contributions of benthic communities to the coverage of the substratum onto the reef slope at 10-m depth (corresponding to fore reef samples n$^2$ 4) are displayed Figure 6. Benthic species were gathered into six categories: mixed algal assemblages, colonizing the limestone and dead corals, coralline algae, *Acropora*, non-*Acropora* corals, and soft corals, as well as abiotic substrate, which integrates sand and rubble. Evaluation of coral categories gives information on structural and functional traits of communities.

Benthic communities were clearly dominated by algal formations that accounted for 59% (BSF) to 79% (SGF) of total community. Hard corals (*Acropora* and non-*Acropora*) represented only from 16% (SGF) to 28% (BSF) of total community (Figure 6). The communities thus appeared to be very homogeneous over the entire outer slope of La Saline reef, in agreement with other regular Global Coral Reef Monitoring Network survey (GCRMN) results. The highest algal coverages were observed south of the Saint Gilles harbour (SGF) and near the Hermitage (HF) and Trois-Bassins passes (SSF, 3BF). Although globally homogeneous from one site to another, the benthic communities differed in the respective composition of the algal formations. Indeed, SGF, TF, PAF, and SSF sites were characterized by a lower proportion of coralline algae (∼3–6%) compared to the other sites (12–23%). In addition, corals were essentially present at TF, PAF, and BSF with *Acropora* mainly found at TF (Figure 6). Non-*Acropora*, dominant at CFF, PAF, and BSF, were mostly represented by shifted coral species: *Asterophora myriophthalma*, *Porites* spp. and *Pocillopora* spp.

**DISCUSSION**

**Biogeochemical Composition of Freshwaters**

The compositions of basalt layer groundwater (BL) and sand layer groundwater (SL) observed here (Figure 2 and Supplementary Figure S1) are in accordance with those reported by Tedetti et al. (2011). They both displayed high Si(OH)$_4$ concentrations as a result of the basaltic bedrock weathering, whereas only BL presented high concentrations of NO$_3^-$ as well as a depletion in DOM and reduced nitrogen. The high NO$_3^-$ content in BL has been proposed to be related to inputs via rainwaters during the transit of groundwater in an agriculture-dominated catchment area. Effectively, groundwater in La Réunion is mainly affected by domestic or livestock effluents (Rogers et al., 2012). On La Saline watershed, NO$_3^-$ contamination was hypothesized to be of agricultural origin (livestock), because NO$_3^-$ infiltration occurred at an altitude where dominant land use activity is livestock farming (Rogers et al., 2012). Fuelled at its base by BL, SL is also fed from the surface by rainwaters and runoffs, allowing the entrance of OM and thus its enrichment in DOM and humic-like material. Moreover, SL presented salinities of 5.7–6.0, suggesting a contamination by marine waters through the rise of the salt wedge during swell events. These marine intrusions in SL may contribute to the decrease in Si(OH)$_4$ and PO$_4^{3-}$ with respect to BL, while the almost complete disappearance of NO$_3^-$ may be presumable due to denitrification (Figure 2 and Supplementary Figure S1). The high content in humic-like material in SL, HR, and 3BR reflected the presence of terrestrial DOM, derived from soil or higher plants (Fellman et al., 2010; Ishii and Boyer, 2012; Tedetti et al., 2012).

Saint Gilles River presented a biogeochemical composition close to that of BL (Figure 2 and Supplementary Figure S1), from which it originates. Indeed, SGR, which is one of the numerous rivers that drain the western flank of the “Piton des Neiges” from 2390 m altitude to the sea, has four basins at different levels from upstream to downstream. These basins are resurgences of BL groundwater escaping from the rock, falling in cascade or going up by the bottom. Although Trois-Bassins River (3BR) is known to be fed by BL (Guigue et al., 2015), its biogeochemical characteristics were much closer to those of SL (Figure 2 and Supplementary Figure S1). 3BR presented low values of Si(OH)$_4$, NO$_3^-$, and PO$_4^{3-}$ very likely due to mixing with marine waters, as shown by the high salinities occurring during swell events. On the other hand, 3BR was enriched in DON, NO$_2^-$, NH$_4^+$, and DOM of terrigenous origin (humic-like) compared to BL and SGR. Its high levels of NH$_4^+$ and NO$_2^-$ are presumably related to the bacterial decomposition of OM.

Hermitage River (HR), which drains SL in the Hermitage sector, presented a high content in terrigenous DOM, as SL and 3BR, and was significantly enriched in DON, tryptophan-like fluorophore and urea compared to all other freshwaters bodies, and was also enriched in PO$_4^{3-}$ compared to SL and the two
other rivers (Figure 2 and Supplementary Figure S1). This enrichment was the sign of anthropogenic pollution through the discharge in the river of a treated wastewater effluent (Tedetti et al., 2011). Actually, the Hermitage wastewater treatment plant, which has a capacity of 25,000 equivalent inhabitants, treats wastewaters from Saint Gilles and Trois-Bassins areas, and releases in HR about 40% of its (tertiary) treated wastewaters. It has been shown that treated and untreated wastewaters contain high amounts of tryptophan-like material (see review by Carstea et al., 2016). Various wastewater organic components could contribute to this tryptophan-like fluorescence signature, such as aromatic amino acids (tryptophan), degradation products of lignin, indoles and aromatic hydrocarbons (Carstea et al., 2016).

**Freshwaters Impacting Reef Waters**

We identified BL groundwater as main source of freshwater, Si(OH)₄ and NO₃⁻ in back reef waters, impacting extensively the southernmost part of the reef (GTB, TEB), and in a lesser extent PAB site, located further north. At these sites, the influence of BL substantially decreased from the beach to the reef flat (Figure 3, Supplementary Figures S2–S5, and Supplementary Tables S3, S4). This is in agreement with previous observations concerning resurgences of BL groundwater in the southern part of the reef (Cuet et al., 1988; Join et al., 1988; Tedetti et al., 2011) or in other reef environments (Paytan et al., 2006; Bowen et al., 2007; Street et al., 2008). Due to the water circulation in the back reef zone, BL groundwater discharged in the southern part of the reef advects northward, parallel to the beach, in a buoyant plume of ~20-m wide (Cuet et al., 2011). The average groundwater discharge along the coast is estimated to be 2.8 m³ d⁻¹ m⁻¹ of shoreline (Join et al., 1988). Humic- and tryptophan-like fluorophores displayed a high spatial variability in the southern part of the reef (Figure 3), reflecting probably in situ biological production in back reef waters due to the NO₃⁻-induced eutrophication, as previously pointed out (Para et al., 2010; Tedetti et al., 2011). Direct inputs from BL are unlikely, as DOM content of BL was low, especially compared to that of SL and HR.

While several works have also reported the evidence of freshwater inputs in the back reef zone (Cuet et al., 2011; Zubia et al., 2018), to our knowledge, the present study is the first one that shows the influence of freshwaters in the outer slope/fore reef waters of La Saline fringing area. Salinity and Si(OH)₄ exhibited a significant inverse correlation for sites PMF, HF, PAF, BSF, SSF, and CFF. This correlation was particularly high for southern sites BSF-3BF and surface waters (Figure 5A and Supplementary Table S5). The salinity-Si(OH)₄ relationship thus represents a first marker of the freshwater discharge in the fore reef waters. These freshwater discharges were more pronounced in the southern stations and surface waters (Supplementary Figures S6B,E), while they were not clearly identified in sites SGE, TF and CFF, neither detected close to the slope at 5–20-m depth (Figure 4B and Supplementary Table S5), as we could have also expected. One possible reason would be that such inputs, if occurring in depth along the slope, would be of very low flowrates in absence of heavy rains (our case) and hence too difficult to catch, either with CTD measurements (Supplementary Figures S5D,E) or with sampling by divers. Although the passes serve as outlets for the waters that have passed through the back reef zone, the freshwater discharges observed here in surface waters at several sites were not only located in front of them. This suggests that additional physical processes were involved in the transfer of waters from the back reef to the fore reef zone, such as swell and the tidal currents (Cuet et al., 2011).

The highest correlations between NO₃⁻, Si(OH)₄ and salinity were observed in the southernmost sites (BSF, SSF, 3BF) (Figure 5B), suggesting a mixture of oceanic waters and nitrate-rich freshwaters. Moreover, the Si(OH)₄/NO₃⁻ ratios in these southern sites were of the same order of magnitude than those recorded in BL (Figure 5B). This confirms that the southern part of the fore reef zone was mainly impacted by BL, which is in accordance with what we observed in the back reef zone (Supplementary Figure S4). Sedimentary formations are poorly developed in this area, and Trois-Bassins River did not contain NO₃⁻ in significant amounts (Figure 2C). Also, the content of humic-like material, which was the main signature of the most superficial freshwaters (SL and rivers, except SGR; Figure 2J), was not very high in the southern part of the reef (Figure 4J). We therefore propose to use the relationships between salinity, Si(OH)₄ and NO₃⁻ to identify and track basalt groundwater discharges in back and fore reef waters (Figures 5A,B and Supplementary Figure S3).

Nevertheless, like NO₂⁻, the spatial distribution of NO₃⁻ was not merely driven by freshwater discharges in surface waters. In fact, many samples with a Si(OH)₄ content close to that of oceanic waters presented a clear enrichment in NO₃⁻ (for instance SFG, PMF, TF, CFF; Figures 4B,C). Regarding the correlations between NO₃⁻, NO₂⁻, and NH₄⁺ (Figures 5C,D and Supplementary Table S5), this NO₃⁻ enrichment seemed to come from an incomplete nitrification of NH₄⁺, derived from another source, or produced in situ by biological communities (Cuet et al., 2011). As a result, the sources of NO₃⁻ in fore reef waters appear multiple (BL groundwater, nitrification...) and may possibly be superimposed. This can contribute to the relative homogeneity of the NO₃⁻ content on the different radials.

The humic-like material reported here, i.e., peaks A + M-component 3 in the Ishii and Boyer (2012)’s classification, has been found in various aquatic environments (Coble, 2007; Fellman et al., 2010; Ishii and Boyer, 2012). It may be of terrestrial origin (derived from soil or higher plants OM) but also come from the transformation of OM formed on the reef, for example in sediments (Burdige et al., 2004; Blanco et al., 2008; Dang et al., 2014). This latter factor may contribute to the overall increase in humic-like concentrations in fore reef waters relative to the oceanic environment, but is unlikely to explain the significant variations observed between sites (Figure 4I). On the other hand, humic-like fluorophore presented the highest concentrations in the most superficial freshwater bodies, except SGR (i.e., SL, HR, and 3BR), while its concentrations in the basalt aquifer were very low (Figure 2J). Consequently, we may probably consider the humic-like material as a good tracer of sand groundwater (SL), and more generally of stagnant and/or superficial freshwaters, such as HR and 3BR. Interestingly,
Kaushal et al. (2020) have shown that this (terrestrial) humic-like material was readily incorporated in coral skeletons, suggesting that coral cores may have the potential to reconstruct past variations in freshwater fluxes.

On the outer slope, humic matter was much more abundant at the surface than at the bottom (Supplementary Figure S7K), suggesting that the contributions occurred in the shallow back reef/reef flat area. Leze (2012) has shown that exchanges between SL and back reef waters occurred in particular due to the rise of the salt wedge during swell events. Indeed, after the rise event, the piezometric level of SL goes down slowly, carrying into the back reef zone waters loaded in compounds (such as humic-like material) present in sedimentary formations (Leze, 2012). Accordingly, the contents in humic-like material were particularly high in sites PMF, HF, and PAF (impacted by SL/HR) compared to the southernmost sites (impacted by BL) (Figure 4J).

Humic-like fluorophore was well correlated to NH$_4^+$ concentration. Since NH$_4^+$ concentrations were quite low in freshwaters, even in SL (~20 μM) (Figure 2E), we may assume that NH$_4^+$ was probably not issued from direct SL or HR inputs. A contribution of NH$_4^+$ produced by reef flat communities (Cuet et al., 2011) is likely in sites where reef water flows back to open sea through passes and outlets. This contribution may be predominant in the southernmost sites (Figure 5E), and especially 3BF where the correlation between humic-like and NH$_4^+$ was relatively low, although significant (Supplementary Table S5). In contrast, high correlations were observed at PMF, HF, and PAF (Figure 5E) suggesting that NH$_4^+$ in fore reef waters may be produced from the photo-oxidation of DOM/humic-like material (Bushaw-Newton and Moran, 1999; Stedmon et al., 2007) or from its bacterial degradation followed by the subsequent excretion activity of grazer organisms (Carlsson et al., 1995). Overall, humic-like material, either issued from direct SL or HR inputs or of reef origin, appeared to be a source of nitrogen to the reef slope. SL or HR inputs of humic-like matter could be particularly significant in the northern part of the reef, where NO$_3^−$ enrichment through BL inputs was not recorded.

Phosphate concentration and tryptophan-like fluorescence exhibited high correlations and the highest values in SGF and HF sites (Figures 4E,F, 5F). This does not seem to be attributable to the biological activity of the reef communities. Although the southern part of the back reef zone was enriched in tryptophan-like (Figure 3G) due to the BL intrusions and subsequent in situ production of OM (Para et al., 2010; Tedetti et al., 2011), the resulting tryptophan export at the Trois-Bassins pass and in the southern fore reef waters was very weak (Figure 4K). In addition, low PO$_4^{3−}$ concentrations were observed at the Trois-Bassins pass (Figure 4F), presumably as a result of this OM production that was accompanied by a consumption of PO$_4^{3−}$ on the reef flat (Cuet et al., 2011).

Within freshwaters, it is only in HR that are found high concentrations of both PO$_4^{3−}$ and tryptophan (Figures 2E,K). As mentioned above, HR is impacted by a treated wastewater effluent and other domestic effluents, which accounts for its high amounts of tryptophan-like material (Carstea et al., 2016). It has been observed significant correlations between tryptophan-like and PO$_4^{3−}$ in river and marine waters affected by wastewater effluents (Baker and Inverarity, 2004; Tedetti et al., 2012). Hence, the correlations found here between tryptophan-like fluorescence and PO$_4^{3−}$ concentration in SGF and HF may be the sign of wastewater pollution, which was confirmed by the analysis of the isotopic composition of nitrate nitrogen in the Hermitage River (Andral et al., 2010). In SGF, this wastewater pollution was probably not issued from SGR, which displayed low contents of tryptophan-like and PO$_4^{3−}$ (Figures 2F,K), but rather from the Saint Gilles harbor or inputs from septic tanks. From these observations, we put forward the use of the relationships between tryptophan-like and PO$_4^{3−}$ to identify and track wastewater inputs in back and fore reef waters (Figure 5F).

Dissolved organic carbon, DON and urea spatial distribution was not influenced by freshwaters discharges in surface waters (Figures 4G–1 and Supplementary Figures S7A–I). Urea concentration in freshwaters was much too low (Figure 2I). While DON concentrations in fore reef waters were in the same range that those in oceanic waters (Figure 4H), DOC concentrations in fore reef waters were even lower than those in oceanic waters, except for BSF2 sample (Figure 4G). Therefore, DOC, DON, and urea did not represent relevant tracers of freshwaters in the reef environment.

### Potential Effects on Phytoplankton Biomass and Benthic Communities

In back reef waters, phytoplankton total biomass was assessed through a proxy commonly used, CTD TChl $a$ concentrations (Supplementary Figure S2), which showed lowest levels north of the Hermitage pass (Figure 3H) where freshwater inputs were not observed (Figure 3A). In fore reef waters, TChl $a$ was opposed to salinity on the second axis of the PCA (Supplementary Figure S8), and the spatial distribution of TChl $a$ (Figure 4L and Supplementary Figure S7R) was therefore roughly reflecting inputs of freshwaters (Figure 4A and Supplementary Figure S6C). These results underline some eutrophication of the southern part of the reef, both in the back reef area as already reported (Cuet et al., 2011; Tedetti et al., 2011; L’Helguen et al., 2014) and in fore reef waters. However, the TChl $a$ distribution pattern did not follow faithfully the distribution pattern of NO$_3^−$ or other nutrients (Figures 4C–F). High TChl $a$ concentrations were not only recorded in the southernmost sites under the direct influence of the NO$_3^−$-rich BL, but also in CMB (back reef) and PAF (fore reef). Conversely, TChl $a$ was very low in HF, while this site was clearly enriched in nutrients due to the influence of the Hermitage River. This discrepancy between the levels of nutrients and TChl $a$ concentrations in HF could reflect an inhibition of photosynthesis by the presence of toxic compounds carried by the Hermitage River, as described through the Aquatic Ecosystem Sustainability Index of toxicity proposed by Barinova (2007). Thus, phytoplankton biomass was not only driven by nutrient inputs in reef waters. Interestingly, we found similar distributions of TChl $a$ and Pheo $a$ concentrations (data not shown for Pheo $a$), with a strong positive correlation between both ($r = 0.92$, $n = 56$, $p < 0.05$). This emphasizes the
strong coupling between phytoplankton biomass and grazing or photo-oxidation processes in fore reef waters.

In the present work, benthic community structure was not determined in the back reef zone. Nonetheless, previous studies showed that all back reef sites, except TB (unaffected by freshwaters), exhibit a decrease in living coral cover, and specifically *Acropora* communities, to the benefit of fleshy algae communities (Mioche and Cuet, 1999; Chazottes et al., 2002; Bruggemann et al., 2008; Naim et al., 2013; Tourrand et al., 2013; Bigot et al., 2016; Zubia et al., 2018; Nicet et al., 2019).

In the same way, over the outer reef slope, a massive presence of algal formations was observed, in particular of mixed algal assemblages, together with a low coral coverage (Figure 6). This trend, i.e., algal formations colonizing bio-constructed substrate and dead corals at the expense of bio-builders (coral species and coralline algae), has been reported in various reef ecosystems (McManus and Polsenberg, 2004; McClanahan et al., 2007; Norström et al., 2009; Bonaldo and Bellwood, 2011). These observations highlight a profound imbalance of the reef environment whatever the outer slope zone considered. This spatio-temporal evolution of benthic communities has been shown for the both sentinel sites of TF and PAF during the 1998–2012 period in the context of the GCRMN survey (Bigot, 2008; data for 2012 from the present study) and confirmed during the last period 2012–2019 (Nicet et al., 2019). Results show that the current trend is part of a regression in the temporal dynamics of corals, explaining the relative homogeneity of benthic community’s health on the outer slope.

Freshwaters, enriched in nutrients and/or OM may reach the fore reef/outer slope environment in two ways. First, they may discharge into the back reef zone before being taken up by the passes (or outlets areas) and dispersed over the outer slope under the effect of tidal currents (the “surface pathway”). Secondly, they may discharge directly on the outer slope at 5–10 m depth through the basalt groundwater complex (the “ground pathway”). Our results indicate that freshwaters mainly reached the outer slope via the surface pathway, while the ground pathway was not really observed (Figure 1). These freshwaters coming from the back reef zone also carried dissolved and particulate OM synthesized on the reef flat, as well as dissolved inorganic nitrogen, in the form of NH$_4^+$, and then affected the benthic communities by diffusion or homogenisation of waters. This explains that high algal coverage rates were more particularly observed around passes, i.e., Hermitage pass (HF site, affected by HR and SL) and north of Trois-Bassins pass (SSF, 3BF sites, affected by BL) (Figure 6). A high algal coverage rate was also found in SFG albeit this site was one of those the less affected by freshwaters (BL and SL). Nonetheless, SFG, like HF, was enriched in tryptophan-like material and PO$_4^{3-}$ attributable to a wastewater pollution. TF and CFF sites, which were not under the influence of freshwaters, also presented significant rates of algal coverage (Figure 6). For several years, high levels of PO$_4^{3-}$ have been detected very regularly on TB reef flat (Cuet, 1989). These have been attributed to a regeneration of dissolved and/or particulate OM mainly issued from the Hermitage pass and secondly transported to TB reef flat by the north tidal current. Tryptophan-like fluorophore has never been measured in parallel, but these high PO$_4^{3-}$ concentrations could be a sign of a wastewater contamination north of the Hermitage pass. In addition, Guigue et al. (2015) have shown that the concentrations in dissolved PAHs were high in TF and CFF sites. Abundant in SSF as well, dissolved PAHs could reflect the presence of sporadic basalt groundwater discharges. The latter could thus affect TF, and to a lesser extent, CFF, in depth outer slope (20–30 m) according to the hypothesis exposed by Join et al. (1988).

It is clear that the freshwater signatures observed mainly in surface waters essentially masked the signal related to the resurgences on the outer slope itself. The resulting nutrient inputs were apparently diffuse, and could be largely assimilated by the algae benthic communities. The potential influence of selected environmental variables on benthic communities in fore reef waters/reef slope was investigated through a CCA (Figure 7). CCA did not reveal any clear influence of environmental parameters on the distribution of corals and mixed algal assemblages. This is not very surprising given the relatively spatial homogeneity of benthic community structure onto the entire slope (Figure 6). However, CCA pointed out an effect of NO$_3^-$, NH$_4^+$, and humic-like fluorophore on the distribution of coralline algae (Figure 7). In other word, the increase in coralline algae abundance may be explained by the increase in NO$_3^-$ and NH$_4^+$ concentrations, and humic-like material. This is rather in accordance with the model of relative dominance proposed by Littler and Littler (1984) in which coralline algae are favored within reef benthic communities in conditions of moderate-to-high N-nutrient concentrations (while high levels of PO$_4^{3-}$ may on the contrary reduce their calcification and growth), coupled to conditions of moderate-to-intense grazing or heavy wave exposure.
CONCLUSION

This study highlights the discharges of basalt (volcanic) groundwater in the southern part, and in a lesser extent, the discharges of sand (sedimentary) groundwater and Hermitage River waters in sites located northward, in both back and fore reef areas of La Saline ecosystem. These discharges mostly impacted fore reef area in surface waters through water transfer from the back reef zone allowed by physical processes (swells, tidal currents) (“surface pathway”). On the other side, basalt groundwater resurgences at 5–20-m depth on the outer slope itself (“ground pathway”) were not clearly detected, due to the diffuse character of these discharges through the basaltic underground/reef transfer, and probably to masking by the freshwater signatures observed in surface waters. Here we propose to use the relationship Si(OH)₄/NO₃⁻ to identify and track basalt groundwater discharges, but also the relationships humic-like fluorescence/NH₄⁺ and tryptophan-like fluorescence/PO₄³⁻ to identify and track sand groundwater discharges and wastewater pollutions, respectively, in reef waters. Besides freshwater discharges, the biogeochemistry of fore reef waters was very likely influenced by autochthonous processes occurring in back reef zone, enhanced or not by freshwater inputs, such as OM bacterial degradation, grazing, photo-oxidation and nitrification. Our results put forward the eutrophication of the southern part of the back and fore reef waters with higher phytoplankton biomasses, even though the latter were not only driven by nutrient inputs. Moreover, a massive presence of algal formations together with a low coral coverage was observed over the outer reef slope, emphasizing the degradation of the reef ecosystem. The spatial homogeneity of the benthic community structures onto the slope contrasted with the spatial heterogeneity of chemical variables in water. This reflects the different spatial and temporal scales involved in biological and chemical processes. In this work, we focused on a summer period for Réunion Island. Future studies taking into account the temporal (seasonal) dimension will undoubtedly help to understand the impact of freshwater discharges and hydrodynamics on the reef biogeochemistry and ecology.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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AUTHOR CONTRIBUTIONS

All authors have participated to the design of the study and sampling/field work and contributed to data treatment. LB and JT were particularly involved in the sampling of water and observation and analysis of benthic communities along the reef slope/in fore reef waters. CG, MG, MT, NF, and PC performed sampling and in situ measurements in freshwaters, back reef and oceanic waters, as well as the analyses of biogeochemical parameters. MT wrote the original manuscript. LB, JT, CG, NF, MG, and PC participated to its writing, reviewing and/or editing.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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