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Habitat suitability and environmental niche comparison of cold-water coral species along the Brazilian continental margin

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ABSTRACT

In face of increasing anthropogenic disturbance in the deep sea, it is a priority to better understand the regional distribution of cold-water corals (CWC). These organisms create some of the most species-rich habitats in the deep sea and, for this reason, they must be properly protected and managed. In this study, we aimed to identify suitable habitat for multiple CWC taxa off the Brazilian continental margin and compare their environmental niches. Habitat suitability models were developed using the Maxent approach, which allowed for the prediction of species distribution and for the identification of potential 'hot spot' areas that may be important for biodiversity conservation. Ecological niches were determined by a PCA-env approach, and niche similarity and equivalence were evaluated based on niche overlap using the Schoener's D metric. Potentially suitable habitat for Octocorallia covered a broad latitudinal range encompassing nearly the entire Brazilian continental margin, whereas Scleractinia had greater potentially suitable habitat in the Central and Southern areas. Scleractinian species were observed to slightly differ in their environmental niche, with non-reef-forming species being more tolerant to a wider range of environmental conditions in comparison with reef-forming species, inhabiting a wider area of the South American continental margin. Due to the high potential suitability for several CWC species, the Central and Southern parts of the Brazilian continental margin should be considered as potential areas high CWC diversity. Considering the current state of the art and strategic assessment tools, these areas are important targets for conservation, management, and environmental impact assessment. Most reef-forming species had similar but not directly equivalent ecological niches, indicating that mapping efforts and management planning should consider CWCs at the species level.

1. Introduction

The deep sea (>200 m depth), the largest biome on Earth that covers 65% of the surface, has been recognized to be environmentally and biologically diverse, providing several important ecosystem services for humans (Thurber et al., 2014). Cold-water corals (CWC) create highly heterogeneous deep-sea habitats that can be used by other organisms as feeding grounds, refuge and/or substrate (Henry et al., 2010; Buhl-Mortensen et al., 2010). In the case of some reef-forming stony coral species (Order Scleractinia) these habitats can be many kilometers long and hundreds of meters high (Roberts et al., 2006). Non reef-forming scleractinian species and the soft corals from the subclass Octocorallia also provide habitat due to their occurrence within dense and often

diverse aggregations known as coral gardens (Roberts et al., 2009) in both hard and soft bottoms. CWC are azooxanthellate corals and present a wide bathymetric distribution generally extending deeper than 50 m water depth (Cairns, 2007). Ecosystems formed by CWC generally have a high biodiversity of associated fauna and are considered some of the most species-rich habitats in the deep sea (Freiwald et al., 2004). However, CWC have a slow growth rate, high fragility (Hall-Spencer et al., 2002) and low recovery potential (Reed et al., 2007; Huvenne et al., 2016), consequently, they are considered to form vulnerable marine ecosystems (VME).

There are six main reef-forming Scleractinian cold-water coral species: Lophelia pertusa (= Desmophyllum pertusum, Linnaeus, 1758); Madrepora oculata Linnaeus, 1758; Solenosmilia variabilis Duncan, 1873;

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Enallopsammia profunda Pourtalès, 1868; *Goniocorella dumosa* Alcock, 1902; and *Oculina varicosa* Le Sueur 1820 (Freiwald et al., 2004). Their importance, distribution and abundance, varies amongst regions (Freiwald et al., 2004). *L. pertusa* is considered as the primary reef-forming species in the deep-sea with an almost world-wide distribution (Davies and Guinotte, 2011). *M. oculata* is widely distributed in the Northeast Atlantic and Mediterranean (Reveillaud et al., 2008; Orejas et al., 2009; Vertino et al., 2010; Gori et al., 2013), whilst *S. variabilis* is concentrated within New Zealand waters (Cairns, 1995; Tracey et al., 2011). In the last decade, habitat suitability models have helped to better understand CWC distribution at both regional and global scales (e.g. Davies et al., 2008; Tittensor et al., 2002; Davies and Guinotte, 2011; Howell et al., 2011; Yesson et al., 2012; Vierod et al., 2014; Georgian et al., 2014; Guinotte and Davies, 2014).

The area predicted using habitat suitability models is generally related to the ecological niche breadth of the species considered (Slatyer et al., 2013). Species that are geographically widespread normally present broader ecological niches given that they persist in an area with wider range of environmental conditions, whereas species with restricted distributions would have a narrower niche breadth (Gaston et al., 1997). Estimating the environmental niche of a given species allows for comparison between different species and may also indicate how sensitive species are to changes in the environment. Although it is possible to test niche similarity from the outputs of habitat suitability models, conceptual and statistical challenges exist with this approach (Broennimann et al., 2012). For example, meaningful niche divergences could be confounded with geographic distance because the environmental data used in species distribution models are often spatially correlated (McCormack et al., 2010). To address this, Broennimann et al. (2012) developed a mathematical approach that can describe species niches in a Principal Component Analyses delimited by the environmental conditions of the study area (PCA-env). This promising tool for conservation and management, allows the evaluation of species niche similarity and serves as a useful complement to habitat suitability models (e.g. Aguirre-Gutiérrez et al., 2015; Zhu et al., 2016).

The exploitation of deep-sea resources is increasing worldwide and there is an urgent need to have an adequate knowledge of the ecology of CWC to ensure that appropriate management strategies are applied (e.g. Morato et al., 2006; Davies et al., 2007). For instance, bottom trawling represents a main threat to CWC (Fosså et al., 2002; Buhl-Mortensen et al., 2016; Buhl-Mortensen, 2017). Oil and gas offshore activities could potentially impact CWC (reviewed in Cordes et al., 2016) and, particularly, the 2010 accidental Deepwater Horizon oil spill in the Gulf of Mexico have showed negative effects on CWC also at 22 km away from the spill site (Fisher et al., 2014) and even 7 years after (Girard and Fisher, 2018). Furthermore, deep-sea mining of massive polymetallic sulfates deposits and cobalt-rich ferromanganese crusts may potentially negatively affect some CWC communities in the near future (Miller et al., 2018).

Three out of six main scleractinian reef-forming species, i.e. L. pertusa, S. variabilis, M. oculata, together with E. rostrata, are known to coexist along the Brazilian continental slope (Castro et al., 2006; Kitahara, 2007; Pires, 2007; Cavalcanti et al., 2017). With a coastline of 7491 km, Brazil accounts for a large proportion of the South American continental margin. This is an extensive area where fisheries, oil and gas exploration are important and ongoing activities. Deep-sea fisheries in the South-eastern Brazilian EEZ (Exclusive Economic Zone) have been estimated to have swept the available area more than once from 2003 to 2011 (Port et al., 2016). In addition, fishing intensity has increased, particularly in the continental slope, from 2000 onwards, due to government policies that stimulated foreign trawlers to occupy deep areas (Perez et al., 2009). Furthermore, the oil and gas production has increased during recent years. Approximately 70% of Brazilian oil production occurs within deep-sea basins (Bernardino and Sumida, 2017). Despite the magnitude and potential environmental impacts, there are few studies analysing the impact of these activities on CWC communities

within this region (e.g. Kitahara, 2009; Port et al., 2016).

In this study, we evaluated the habitat suitability distribution and niche of several CWC species along the Brazilian continental margin and slope. We aimed to better understand differences in CWC potential distribution and corresponding environmental niches. We compared the Octocorallia Subclass and Scleractinia Order and then focused on several scleractinian species. Six species of scleractinians were studied in order to compare their niche and potential distributions representing diverse types of CWC VMEs. Four reef-forming species L. pertusa, M. oculata, S. variabilis and E. rostrata and two non-reef-forming corals, Cladocora debilis Milne Edwards and Haime, 1849 and Deltocyathus Milne Edwards and Haime, 1848. The former type of species represents reef presence or possible reef-habitat formations and the latter type represents possible coral fields, being C. debilis a colonial species and Deltocyathus spp. a solitary cup-coral genus. The results presented here are a fundamental step in better understanding CWC distribution and their possible sensitivity to environmental changes both natural and human-induced within the Brazilian continental margin and slope.

2. Methods

2.1. Study area

The present study focused on the Brazilian continental margin and slope, but also included international waters to 25° W (Fig. 1 a). The Brazilian continental margin was divided into four areas from north to south based upon their oceanographic and biological characteristics (after the REVIZEE program - Brazilian project on Living Resources in the Exclusive Economic Zone, Anon, 2006, www.mma.gov.br/revizee) as follows: (1) The North Brazil Shelf including the Amazonia, hereafter referred as the "North Area"; (2) the tropical north-eastern Brazil area, including São Pedro and São Paulo and Fernando de Noronha archipelagos and Rocas Atoll as the "North-eastern Area"; (3) the Eastern Brazil and Trindade and Martin Vaz Islands as the "Central Area "; and (4) the warm temperate South-eastern Brazilian area as the "South Area ", including the Rio Grande Rise (Fig. 1 b).

2.2. Species data

A database was compiled using all available CWC records in the Brazilian continental margin from three sources: (1) records in available



Fig. 1. (a) Overview of the study area localization. (b) The Brazilian continental margin with the CWC presence points of Octocorallia and Scleractinia (light blue and light grey, respectively), in the focal areas used in the present study. The boundaries of the Brazilian EEZ along the Brazilian coast are represented by the grey line. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

databases, including the Ocean Biogeographic Information System (OBIS) and the REVIZEE program; (2) records from specific literature sources, and (3) field observations using remotely operated vehicles in Campos Basin (Cavalcanti et al., 2017) (Table 1, Supplemental material). Octocorallia records were mainly from the North and Central Brazilian areas and principally from 30 to 150 m depth, i.e. mesophotic habitats. In order to exclude zooxanthellate corals, we used only records deeper than 50 m (Cairns, 2007). To model habitat suitability of each taxon, all records were filtered to provide only one record per analysis cell (ca. 1 km²) (Table 1, Fig. 1b). We selected mostly published species records to ensure a good taxonomic classification at species level. Furthermore, records from other sources were used at taxonomic level higher than genera to avoid possible identification errors.

Table 1

Number of occurrence records of azooxanthellate corals from the Brazilian continental margin and slope, including historical records from published sources (see references in supplementary Table 1) and new scleractinian records from Cavalcanti et al. (2017), and the filtered total number of records (with one record for analysis cell) that were used to model habitat suitability.

Taxonomic group	Historical records	New records	Total records used in models
Order Scleractinia	396	1147	259
Subclass	151		60
Octocorallia			
Species			
Cladocora debilis	57		54
Deltocyathus spp.	36		33
Enallopsammia	8	222	33
rostrata			
Lophelia pertusa	77	342	75
Madrepora oculata	21	97	29
Solenosmilia variabilis	26	486	72

2.3. Environmental data

A total of 34 environmental variables were used to model species habitat suitability (after Davies and Guinotte, 2011). They were classified in seven broad categories (after Yesson et al., 2012): carbonate variables (CARB), bathymetric variables (BATH), hydrodynamic variables (HYDRO), productivity variables (PROD), oxygen variables (OXY), chemical variables (CHEM), and temperature variable (TEMP) (Table 2). These categories were selected based on biological knowledge or single-factor analysis (Yesson et al., 2012). After subset the variables into top-level categories, only a single variable per category was selected by creating species distribution models for each variable on its own, calculating the validation area under the curve (AUC, see below) and retaining the most important variable per category for the final analysis (Phillips et al., 2009).

2.4. Maxent model predictions

Maxent version 3.3.3k (Phillips et al., 2006) was used to model habitat suitability. Presence records were randomly divided in two subsets, one with 70% of records to train the model and one with the remaining 30% to testing the models. Models were run with the default parameters of Maxent i.e., convergence threshold of 10^{-5} , a maximum of 500 iterations, and a regularization multiplier of 1. The importance of each variable on each final model was assessed using a Jackknifing procedure (Phillips et al., 2009). This procedure compares the contribution of each variable between two models, one without the variable and a second including it, thereby determining how much new information the variable contributes to a model and how much is lost when that variable is omitted. Final model predictions were presented as maps showing a predicted continuous habitat suitability value. In order to contrast predictions from different taxonomic groups, these outputs

Table 2

Environmental variables used to the variables selection divided in seven categories: carbonate variables (CARB), bathymetric variables (BATH), hydrodynamic variables (HYDRO), productivity variables (PROD), oxygen variables (OXY), chemical variables (CHEM), and temperature (TEMP).

	VARIABLE	NAME	REFERENCE	UNIT
CARB	Aragonite saturation state	arag_orr	Orr et al. (2005)	Ω_{ARAG}
	Aragonite saturation state	arag_stein	Steinacher et al. (2009)	Ω_{ARAG}
	Calcite saturation state	cal_orr	Orr et al. (2005)	Ω_{CALC}
	Calcite saturation state	cal_stein	Steinacher et al. (2009)	Ω_{CALC}
BATH	Aspect	aspect	Jennes (2013)	degrees
	Aspect- Eastness	eastness	Wilson et al. (2007)	-
	Aspect- Northness	northeness	Wilson et al. (2007)	-
	Curvature - Plan	plancurve	Jennes (2013)	-
	Curvature - Profile	profilecurve	Jennes (2013)	-
	Curvature - Tangential	tangcurv	Jennes (2013)	-
	Roughness	roughness	Wilson et al. (2007)	-
	Rugosity	rugosity	Jennes (2013)	-
	Slope	slope	Jennes (2013)	degrees
	Bathymetry	srtm30	Becker et al. (2009)	m
	Terrain Ruggedness Index	tpi	Wilson et al. (2007)	-
	Topographic Position Index	tri	Wilson et al. (2007)	-
HYDRO	Current velocity	regfl	Carton et al. (2005)	$m s^{-1}$
	Vertical current velocity	verfl	Carton et al. (2005)	$m s^{-1}$
PROD	Seasonal variation index	lutzs	Lutzs et al. (2007)	-
	Primary productivity (maximum)	modismax	NASA Ocean Color	$g C m^{-2} y^{-1}$
	Primary productivity (mean)	modismean	NASA Ocean Color	$g C m^{-2} y^{-1}$
	Primary productivity (minimun)	modismin	NASA Ocean Color	$g C m^{-2} y^{-1}$
	Particulate Organic carbon	poc	Lutz et al. (2007)	$g C_{org} m^{-2} y^1$
	Primary Productivity (Vertically generalised	vgpmean	Behrenfeld and Falkowski (1997)	$g C m^{-2} y^{-1}$
	productivity model)			
OXY	Apparent oxygen utilisation	oaxu	García-Alegre et al. (2014)	mol O ₂
	Percent oxygen saturation	poxs	García-Alegre et al. (2014)	% O ₂
	Dissolved oxygen	disso2	García-Alegre et al. (2014)	$ml l^{-1}$
CHEM	Nitrate	nit	García-Alegre et al. (2014)	μ mol l ⁻¹
	Phosphate	phos	García-Alegre et al. (2014)	μ mol l $^{-1}$
	Salinity	sal	Boyer et al. (2005)	PSS
	Silicate	sil	García-Alegre et al. (2014)	μ mol l $^{-1}$
TEMP	Temperature	temp	Boyer et al. (2005)	°C

were converted into binary values (0 = unsuitable, 1 = suitable) based on the maximum sum of sensitivity plus specificity test threshold (Jiménez-Valverde and Lobo, 2007; Liu et al., 2013). This representation allowed to identify potentially suitable areas for multiple taxon as potential areas of hotspot of cold-water coral diversity.

2.5. Species niche comparisons

In order to describe and compare the environmental niches, we have represented the species niche in a multivariate space, measured the niche overlap and tested their similarity and equivalence. The spatial niche occupied by each species was represented by an environmental principal component analysis (PCA-env) approach (after Broennimann et al., 2012) using the seven most important environmental variables for each taxon (see environmental data session for the variables selection). Despite differences between Octocorallia and Scleractinia carbonate structures (calcite vs aragonite, respectively), a unique environmental space is needed to compare their species niche. Then, whilst not biologically relevant for Octocorallia, aragonite saturation state (AST), which was highly correlated with the Calcite saturation state (CST) (r > 0.9), was used to construct the PCA-env.

Species occurrences were disaggregated with a minimum distance equal to the grid resolution (ca. 1 km) to reduce sampling bias. The environmental space, delimited by the axes, was set to 100×100 cells. These cells were used to generate a 'smoothed' Kernel density of the filtered occurrences, as well as of the available environmental conditions (in the focal areas, Fig. 1). The occupancy of the environment by the entity was calculated in each cell (Broennimann et al., 2012). Niche overlap was estimated using Schoener's D metric, which ranges from 0 (no overlap) to 1 (complete overlap) (Schoener, 1970). This metric is used to test niche similarity and equivalence, comparing the observed D value and the probability distribution of overlap values resulted from simulated species occurrence distributions (Warren et al., 2008). The application of a smoother kernel to standardize species densities allowed moving from the geographical space to the multivariate environmental space, making the analysis independent of sampling effort and environmental space resolution (Broennimann et al., 2012).

Similarity and equivalence tests are used to evaluate if the environmental niches of two species are more similar than expected by chance and if both species have the same use of the niche space, respectively (Warren et al., 2008). The similarity test for each species pair (a and b) consisted of two reciprocal comparisons. The observed D value is compared with the probability distribution of overlap values created from simulations of random points in the background area. Then, the real overlap is compared with the overlap between randomly distributed species in the area (same n of species presence points and with "background" being the studied area). Whereas equivalence test consisted in the comparison of the observed D value and the probability distribution of overlap values from simulated species distribution randomly created from the real presence points of both species (Warren et al., 2008). Since the species have broad regional and global distributions, high dispersal capacity is expected, thus, the background area represented a variety of ecological space that they could colonize as suggested by Barve et al. (2011). The tests were based on 100 iterations. All the analyses were done in R (R Development Core Team, 2010) with the "ecospat" package (Broennimann et al., 2016).

Niche density center and niche breadth were calculated to determine how the niches differed, considering the two main principal components. The former indicates the optimal environmental conditions for the species, while the latter described the capacity of a species to tolerate deviations from the optima. Smaller values indicated that the species occupies a more restricted or specialized niche, and larger values that the species has a wider environmental niche. Niche density center was calculated as the mean of the Kernel's density value calculated from 10000 random points inside of the environmental niche space of each taxon. Niche breadth was estimated as the proportion of the available environmental conditions delimited by the axes $(100 \times 100 \text{ cells})$ that were estimated as occupied in the PCA-env, i.e., the percentage of available conditions inhabited by the species (represented as percentage). To better represent niche center position differences, the change from Scleractinia niche center to the specific species was represented with an arrow in the representation of its environmental niche plot.

3. Results

3.1. Occupied environmental conditions

The environmental conditions occupied by Octocorallia and Scleractinia were mostly in waters saturated with calcite and aragonite respectively (i.e., $\Omega \ge 1$), although there were some Scleractinia occurrences found in aragonite-depleted waters (Fig. 2a). Octocorallia and Scleractinia were mainly present in depth shallower than 1250 m (Fig. 2b). Octocorallia occurrences were found across a broad range of current velocities from 0 to 0.20 m s⁻¹, whereas Scleractinia were found at sites with velocities lower than 0.1 m s⁻¹ (Fig. 2c). Both, Octocorallia and Scleractinia were found in sites with POC that ranged from 0 to 50 gC C_{org} m⁻² y⁻¹ and in a range of dissolved oxygen conditions ranging between 4 and 5.5-ml l⁻¹, and silicate <50 µmol l⁻¹ (Fig. 2d, e and f, respectively). Octocorallia occupied a wider temperature range (ca. 0–30 °C), while Scleractinia occupied waters less than 25 °C (Fig. 2g).

The individual scleractinian species had slight differences between them: *E. rostrata*, *M. oculata*, *L. pertusa* and *S. variabilis* were restricted to current velocities lower than 0.5m s^{-1} (Fig. 2 j) and, together with *C. debilis* were limited to sites with POC lower than 20 g $C_{org} \text{ m}^{-2} \text{ y}^{-1}$ (Fig 2 k); *C. debilis* and *Deltocyathus* spp. were found in a smaller range of dissolved oxygen conditions, between 4.5- and 5.5-ml l⁻¹ (Fig 2 l), and were found mainly in waters with silicate concentrations < 20 µmol l⁻¹ (Fig. 2 m); *S. variabilis* had the narrowest temperature range, with most occurrences in a maximum of 10 °C (Fig. 2 n).

3.2. Variable selection and contribution

Variables with the highest AUC from each category were selected for model construction, with the exception of aragonite saturation state (AST), which was selected for Scleractinia and each scleractinian species as they have aragonitic skeletons. Variables selected for final models of Octocorallia and Scleractinia were the same, with the exception of the carbonate variable (Table 3). The variables that most contributed to the final model for Octocorallia were calcite saturation state (CST), dissolved oxygen and temperature, whereas for Scleractinia were AST, depth and temperature (Table 4). Variables selected for specific models of Deltocyathus spp., E. rostrata, M. oculata and L. pertusa were AST, depth, current velocity, POC, dissolved oxygen, silicate and temperature (Table 3). Whereas for C. debilis oxygen utilisation was selected instead of dissolved oxygen and for S. variabilis salinity in place of silicate (Table 3). The final models demonstrated differences in the three variables that contributed most. C. debilis, Deltocyathus spp., E. rostrata, L. pertusa and M. oculata all showed that AST, depth and temperature were the best contributors for these species, whilst depth, POC and salinity were best for S. variabilis (Table 4).

3.3. Model performance and habitat suitable area

Octocorallia and Scleractinia models performed well, with AUC of 0.9681 and 0.9551 from test data, respectively. The distribution of suitable habitat for Octocorallia encompassed along the entire Brazilian continental margin, including the Vitória-Trindade seamount chain and Rio Grande Rise (30°S, 35°W) (Fig. 3). Whereas the habitat suitability distribution of Scleractinia was more restricted to the Central and South area of Brazil, between 20°S 37°W and 30°S 50°W, including the Rio Grande Rise (Fig. 3, ESM Figs. 1 and 2, respectively). Both, Scleractinia and Octocorallia demonstrated an overlap in suitable habitat within the



Fig. 2. Kernel density curves of the environmental conditions occupied by corals from the Brazilian continental margin. Octocorallia (continuous line) and Scleractinia (dotted line) are shown to the left of the figure, and *Cladocora debilis* (black dotted line), *Deltocyathus* spp. (yellow dotted line), *Enallopsammia rostrata* (red line), *Lophelia pertusa* (dark green line), *Madrepora oculata* (grey line), and *Solenosmilia variabilis* (blue line), to the right. Carbonate condition represent aragonite saturation state (Ω) for Scleractinia and scleractinian species, and calcite saturation state for Octocorallia. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

Central and South region, as well as in the Rio Grande Rise (Fig. 3).

The scleractinian species-level models also exhibited acceptable model performance (Test-AUC: *C. debilis*, 0.9857; *Deltocyathus* spp., 0.9412; *E. rostrata*, 0.997; *L. pertusa*, 0.9902; *M. oculata*, 0.975; *S. variabilis*, 0.9935). Both, *C. debilis* and *Deltocyathus* spp., demonstrated suitable habitat around the South and Central areas, between 20°S 40°W and 42°S 23°W, covering a greater area of continental slope, in terms of bathymetric extent, than reef-forming species (ESM Figs. 3 and 4). The reef-forming species *L. pertusa* and *M. oculata* were concentrated around the Northeast, Central and South area (ESM Fig 5 and 6), whilst *E. rostrata* and *S. variabilis* were restricted to the Central and South area, especially the Vitória-Trindade seamount chain and Campos and Santos Basins (i.e., between 20°S 40°W and 42°S 23°W) (Fig. 3 B and ESM Figs. 7 and 8). Most scleractinian species overlapped in predicted suitable habitat in some regions of the Northeast area (Fig. 3 b, i and ii); the Central area (in Vitoria-Trindade chain) (Fig. 3 b, ii); the South area (Fig. 3 b, iv); and the Rio Grande rise (Fig. 3 b, iii).

3.4. Ecological species niche comparison

Environmental niches of Scleractinia and Octocorallia were determined mainly by temperature, depth, AST, POC and regional flux (water current velocity) on the principal component 1 (PC1) and by dissolved oxygen and silicate on the second principal component (PC2). These two components explained ca. 65.5% and 18.5%, respectively, of the environmental conditions occupied (Fig. 4 c). Octocorals had a wider environmental niche than Scleractinia, mainly along the PC1, covering 30% and 16% of the environmental conditions of the region, respectively (Fig. 4 a and 4 b). The center of density of niches were highly differentiated between both groups reflecting the low niche overlap between

Table 3

Test AUC values for Maxent model of Octocorallia and Scleractinia taxa and for six scleractinian species in the Brazilian continental margin, based in a single variable. A value close to 0.5 indicates a model no better than a random prediction, values greater than this and closer to 1 indicate models with better predictive power. A value of 1 indicates a theoretically perfect model. Values in bold indicate the main variable of each categorical group and which were selected to run the final models with the exception of arag_orr which was used to Scleractinia and the scleractinian species due their ecological importance (See section 2.2. Variables selection and contribution). Category and variable name abbreviations are presented in Table 2.

	Variable	Scleractinia	Octocorallia	C. debilis	Deltocyathus spp.	E. rostrata	L. pertusa	M. oculata	S. variabilis
CARB	arag_orr	0.9525	0.9248	0.9671	0.953	0.9612	0.9624	0.9586	0.9604
	arag_stein	0.9562	0.9233	0.9691	0.9234	0.9588	0.9427	0.9757	0.9135
	cal_orr	0.9579	0.9269	0.9663	0.9524	0.9648	0.9634	0.9637	0.9736
	cal_stein	0.9567	0.9206	0.9697	0.9255	0.9642	0.9487	0.9763	0.9256
BATH	srtm30	0.9664	0.9311	0.9737	0.9659	0.9913	0.9877	0.9951	0.9885
	roughness	0.5204	0.5195	0.475	0.6183	0.6258	0.5883	0.5	0.6505
	rugosity	0.5518	0.6296	0.5243	0.6131	0.3548	0.5759	0.3915	0.6288
	slope	0.599	0.7215	0.4622	0.5222	0.7734	0.7702	0.8024	0.756
	TPI	0.7852	0.5012	0.7461	0.782	0.8196	0.7119	0.9248	0.8564
	TRI	0.7088	0.483	0.6351	0.5874	0.7472	0.6516	0.8489	0.7573
	aspect	0.7332	0.4834	0.6824	0.7362	0.7748	0.6292	0.8729	0.744
	eastness	0.4767	0.5079	0.4937	0.5686	0.431	0.6141	0.5	0.6238
	northeness	0.5467	0.6074	0.5552	0.6484	0.4031	0.5142	0.5	0.5696
	plancurve	0.602	0.7327	0.4447	0.5135	0.763	0.7649	0.8147	0.7787
	longcurve	0.7057	0.7277	0.4079	0.5455	0.7977	0.7677	0.8069	0.7629
	tangcurve	0.5981	0.7435	0.4055	0.5352	0.7673	0.7676	0.8106	0.7608
HYDRO	regfl	0.8705	0.9231	0.9187	0.7648	0.9422	0.8762	0.9566	0.8705
	verfl	0.7004	0.6403	0.33	0.2592	0.743	0.2374	0.7354	0.5
PROD	lutzs	0.8487	0.7848	0.947	0.9316	0.9044	0.8349	0.9645	0.7948
	modismax	0.8086	0.8456	0.9491	0.7766	0.9128	0.8689	0.9669	0.8357
	modismean	0.8351	0.8334	0.9498	0.8157	0.9265	0.9004	0.9779	0.8681
	modismin	0.8086	0.8456	0.9491	0.7766	0.9128	0.8689	0.9669	0.8357
	POC	0.9369	0.8621	0.9662	0.9555	0.9878	0.9727	0.9931	0.978
	vgpmean	0.8439	0.809	0.9544	0.8389	0.9449	0.8847	0.9706	0.8819
OXY	aoxu	0.8195	0.8954	0.9126	0.677	0.6107	0.7846	0.6575	0.6625
	disso2	0.8913	0.939	0.8654	0.8969	0.9083	0.9375	0.9671	0.9477
	poxs	0.7796	0.8992	0.894	0.6299	0.5998	0.7486	0.4196	0.7295
CHEM	dic_stein	0.8889	0.9214	0.9345	0.7986	0.8179	0.7632	0.9426	0.6993
	nit	0.8281	0.8829	0.8761	0.7474	0.5964	0.8172	0.6385	0.6625
	phos	0.8442	0.9249	0.8669	0.8148	0.6673	0.868	0.6667	0.7377
	sal	0.8962	0.9082	0.8425	0.6082	0.784	0.8779	0.5401	0.9445
	sil	0.9145	0.9273	0.9422	0.8868	0.9312	0.9158	0.9782	0.8753
TEMP	temp	0.9618	0.9309	0.9678	0.953	0.9855	0.9825	0.9951	0.9776

Table 4

Test AUC values for Scleractinia, Octocorallia and six different scleractinian species models, based in a single variable model. Values corresponding to the three most significant variables for each taxon are in bold. Variable name abbreviations are presented in Table 2.

	Scleractinia	Octocorallia	C. debilis	L. pertusa	M. oculata	S. variabilis	E. rostrata	Deltocyathus spp.
AUC								
calc-orr		0.9403						
arag-orr	0.9457		0.9694	0.9565	0.9543	0.9671	0.9403	0.9683
srtm30	0.9518	0.9492	0.9784	0.9882	0.9812	0.9877	0.9493	0.9595
regfl	0.8812	0.8234	0.9223	0.8743	0.916	0.8874	0.8234	0.6203
POC	0.9214	0.8326	0.9685	0.9335	0.8644	0.9748	0.8326	0.9434
diso2	0.8941	0.9206		0.934	0.8869	0.9431	0.9206	0.8913
aoxu			0.8877					
sil	0.8864	0.8639	0.9358	0.9088	0.9527		0.8639	0.9484
sal						0.9905		
temp	0.9424	0.9281	0.9698	0.98	0.9761	0.969	0.9281	0.9635

Octocorallia and Scleractinia (D = 0.28) (Table 5).

The environmental niche of Scleractinia presented two subcenters; corresponding to reef-forming and non-reef-forming species (Fig. 4 b). The non-reef-forming species, i.e. *C. debilis* and *Deltocyathus* spp., had their density center displaced to the down-left indicating that their niches were determined by higher temperature, POC, current velocity, AST and dissolved oxygen, as well as shallower depths and lower silicate concentration (Fig. 4 d and e). Both species had an environmental niche that covered the 8% of the environmental conditions available in the region. Reef-forming species had the niche density center displaced to the top-right showing opposite patterns to non-reef-forming species (Fig. 4 f–i). The environmental niche of *E. rostrata, L. pertusa, M. oculata* and *S. variabilis* covered 4%, 8%, 7% and 5% of the environmental conditions available in the region, respectively. The similarity niche

hypothesis was accepted for most pairs of scleractinian species in both directions, except for *E. rostrata* compared to *C. debilis*, *Deltocyathus* spp. and *L. pertusa*, and for *M. oculata* compared to *S. variabilis*, in both directions (Table 5). The niche equivalence hypothesis was rejected for all Scleractinia species.

4. Discussion

In this study, we developed habitat suitability distribution models built specifically for the main CWC taxa found along the Brazilian continental margin. These results represent significant improvement over previous model predictions for this region, which were derived from global scale predictions (i.e. Davies et al., 2008, 2011; Yesson et al., 2012). This improvement is largely determined by the new



Fig. 3. Potential distribution areas and the overlap between the different taxa. A) Octocorallia (light blue) and Scleractinia group (light grey) with their overlapped area (marine blue). Main overlapping areas: Central area, Rio Grande rise, South area; B) Scleractinian species (*Cladocora debilis, Deltocyathus* spp., *Enallopsammia rostrata, Lophelia pertusa, Madrepora oculata* and *Solenosmilia variabilis*) overlap, showed as the number of species potentially distributed in the same area, along the Brazilian continental margin. Areas with higher number of species potentially distributed there: i) north of the Northeast score in front of the Rio Grande do Norte state, ii) east slope of the Northeast area in front of Bahia state, Central area, including the margins of submersed islands of the Vitoria-Trindade chain, iii) Rio Grande rise, in Southwestern Atlantic International waters, and iv) South area, from Cape of São Tomé to 30° S in Rio Grande do Sul state. Potential distribution areas are represented with the maximum sensitivity plus specificity test threshold. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

region-specific compilation of occurrence records and also the construction of a regional scale model, which has a higher capability to identify specific niche features as result of the use of a more focused range of background environmental conditions (Vierod et al., 2014). In addition, widely distributed species could exhibit regional or local niche differences for a variety of evolutionary and non-evolutionary reasons (Lesica and Allendorf, 1995; Leibold et al., 2019) that are not captured in larger scale models. Representing regional ecological and environmental features allows for a stronger local predictive power (Osborne and Suarez-Seoane, 2002; Murphy and Lovett-Doust, 2007). This approach can thus greatly enhance the understanding of CWC habitat suitability within the Brazilian continental margin and also allowed for an evaluation of the overlap between the CWCs found within the region.

We observed geographical overlap in the predicted suitable habitat for Octocorallia and Scleractinia, as well as between individual scleractinian species in the Campos Basin, Santos Basin, Vitória-Trindade chain, Rio Grande rise and Pelotas Basin (South area). As CWCs constitute high biodiversity habitats in the deep (Henry and Roberts, 2017), we suggest that these areas may be of special interest, particularly for biodiversity conservation. This reinforces previous studies that highlight the regional importance of the Vitória-Trindade chain as a priority area for conservation (Meirelles et al., 2015; Pinheiro et al., 2015) and addresses the lack of information about the deep-sea biodiversity in the region. In the Southern area and Rio Grande Rise, however, the substantial overlap in Octocorallia and Scleractinia habitat suitability needs to be validated given there are no published presence records of octocorals from the Southern area, and there are no CWC records from the Rio Grande Rise. This highlights the need of surveys in these regions, particularly in Rio Grande rise, which could be a site for future mineral extraction of cobalt crusts, which could impact coral communities and their associated biodiversity if they are found there.

that encompassed nearly the entire Brazilian continental margin, with large areas of highly suitable habitat in the Central, North and Northeast areas. In the North area (Amazon Basin), there was high predicted suitability for Octocorallia in the mesophotic region, which agrees with the many occurrence records (approximately 75%) that were between 50 and 150 m depth. Their distribution around this region has been recently corroborated by the observation of several important communities (Cordeiro et al., 2015; Moura et al., 2016). Even though there are no published records of octocorals in the South area and the Rio Grande Rise, where our models have shown potentially suitable conditions (Fig. 2), several unpublished records from these areas validate our results (M.V. Kitahara and P.Y.G. Sumida pers. obs.). The wide distribution of Octocorallia was related with a wide environmental niche breadth, which covered 30% of the available environmental conditions of the Brazilian continental margin, suggesting that they may be more common than previously thought.

In contrast, Scleractinia had a far narrower habitat suitability distribution, mainly along the Central and South areas of the Brazilian continental margin, and a lower habitat suitability around the Northeast. The deep sea in these three areas is dominated by the presence of Antarctic Intermediate Water (AIW), which has been found to be associated with several scleractinian species in this region between 550 and 1200 m depth (Viana et al., 1998; Sumida et al., 2004; Arantes et al., 2009). The broader habitat suitability distribution of Octocorallia compared to Scleractinia can be translated into a potentially higher tolerance to physical-chemical conditions in the environment by Octocorallia (e.g. mesophotic corals at shallower depths), which generally allows for a wider geographic distribution (Slatyer et al., 2013).

Substrate is a fundamental factor for the settlement of CWC and its availability strongly control their colonization. Nevertheless, the availability of this information is generally limited, particularly for big areas as the entire Brazilian continental margin. Therefore, the realized

Suitable habitat for Octocorallia covered a broad latitudinal range



Fig. 4. Ecological niche of the cold-water coral species studied in the environmental space produced by the principal component analysis method and the variables contribution for loading their two principal components (PC) (c). The results represent the niche of the species in the two main axes determining different environmental condition. The grey to black shading represents the grid cell Kernel density of the species occurrences, black being the highest density. Dashed lines represent the 50% of the available environmental conditions in the Brazilian continental margin, and the solid line represent the 100%. Red arrows connect density center of the species niche to the density center of Scleractinia and illustrate their differences. In (c), POC in abbreviation for particulate organic carbon and AST, aragonite saturation state. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

distribution of corals could be far smaller in area than the potential one we are presenting here (Guinotte and Davies, 2014; Anderson et al., 2016). CWC that settle in hard-bottom substrate are likely more restricted than soft-bottom substrate colonizing species. Then, the larger morphological variety of octocorals, which can present a peduncle for anchoring in mud and sand (order Pennatulacea), a basal disk for fixation in hard substrates, or a basal rootlike processes to anchor in sediment substrates (Bayer, 1961), may allow then to inhabit both substrate types. Contrary to Octocorallia, most scleractinians are restricted to hard-substrate, but some species, such as the cup coral Deltocyathus genus inhabit soft substrates. For instance, octocorals have been observed colonizing shells and rock fragments due to the scarcity of common hard substrates in the Amazon Basin (Cordeiro et al., 2015). Whereas in the southeastern area (in Campos basin), the occurrence of diverse Octocorallia species indicated the presence of both hard and soft habitats along the continental slope (Arantes et al., 2009). Arantes et al. (2009) observed co-occurrence of octocorals of both type of substrates and scleractinians in the middle slope, which corroborates the observed overlap in their suitable distribution (Fig. 3 a-i).

CST, depth, and temperature accounted for the highest contributions to corals suitability predictions and agree with findings from previous studies about CWC distributions (Davies et al., 2008; Dullo et al., 2008;

Tittensor et al., 2009; Davies and Guinotte, 2011). CST and AST have been found to be strong predictors in many CWC species distribution models, mainly at global scales (Davies et al., 2008; Tittensor et al., 2009; Davies and Guinotte, 2011; Yesson et al., 2012) and in some specific regions, such as the Pacific Ocean (Anderson et al., 2016). POC is the main food source for many cold-water coral species (Kiriakoulakis et al., 2004, 2007), but in this study it was an important predictor only for S. variabilis. POC concentration was relatively low in many areas where corals were found, as has been observed in other regions, such as in the Northwestern Pacific (Guinotte and Davies, 2014). Nevertheless, these low values observed here could be due to the fact that POC was calculated as an annual mean and would underrepresent important seasonal pulses. For instance, increases in POC resulting from down-welling events or water movements due to internal waves are important for some coral species, such as L. pertusa (Davies et al., 2009; Purser et al., 2010; van Oevelen et al., 2016). Silicate environmental conditions have been indicated to have a negative relationship with L. pertusa global distribution (Davies et al., 2008), and a strong negative correlation with coral species richness (especially within the north-east Pacific) (Reyes Bonilla and Cruz Piñón, 2002). Low silicate concentration is associated with low primary productivity waters (Longhurst and Pauly, 1987). Here, silicate was an important factor to predict species

Table 5

Niche comparison for cold water corals in the Brazilian continental margin. Niche overlap values Schoener's D (Schoener, 1970; Broennimann et al., 2012) and the significance of similarity and equivalence tests are giving for each pair-wise comparison (ns: not significant). The higher overlap value is presented in bold and the lowest in italics.

Táxon		Niche Overlap (D)	Niche similarity	Niche equivalence
а	b		a – b and b -	
			а	
C. debilis	Deltocyathus	0.89	Similar	Different
	spp.			
	E. rostrata	0.01	ns	Different
	L. pertusa	0.72	Similar	Different
	M. oculata	0.32	Similar	Different
	S. variabilis	0.008	Similar	Different
Deltocyathus	E. rostrata	0.04	ns	Different
spp.				
	L. pertusa	0.80	Similar	Different
	M. oculata	0.40	Similar	Different
	S. variabilis	0.035	Similar	Different
E. rostrata	L. pertusa	0.21	ns	Different
	M. oculata	0.48	Similar	Different
	S. variabilis	0.77	Similar	Different
L. pertusa	M. oculata	0.57	Similar	Different
	S. variabilis	0.20	Similar	Different
M. oculata	S. variabilis	0.41	ns	Different
Octocorallia	Scleractinia	0.28	ns	Different

suitability distribution. Silicate was related with depth and could be an indicator of productivity along the Brazilian continental margin that particularly differentiated the more productive conditions inhabited by Octocorallia compared with Scleractinia.

There were clear differences in environmental niche of the scleractinian species investigated in this study. Reef-forming species were found in conditions with higher silicate and depth but in a lower dissolved oxygen concentration range (a minimum of 4.5 ml l^{-1}), AST, POC, temperature and regional flow conditions compared with non-reefforming species. Non-reef-forming species niches were associated to a larger range of silicate conditions, regional flow and POC conditions (Figs. 2 and 4). These results suggest that non-reef-forming species are not limited by environmental conditions as the studied reef-forming species within the Brazilian continental margin. This is corroborated by their wide distribution in the neighbor Colombian Caribbean region between 10-153m and 70–520m, respectively (Santodomingo et al., 2013).

Based on the niche conservation theory ecological niches are thought to be more similar between close-related taxa (reviewed in Wiens and Grahan, 2005). Within both groups, reef forming and non-reef-forming species, there were significant niche similarities and differences that were not related with their phylogenetic proximity. For instance, E. rostrata and M. oculata are two reef-forming species that present similar environmental niches and belong to different clades of Scleractinia, i.e., the "Complex" and "Robust" clades, respectively (Stolarski et al., 2011). On the other hand, E. rostrata and Deltocyathus spp., a reef-forming and non-reef-forming species, respectively, are from the "Complex" clade but presented a non-similar environmental niche. This suggests that the differentiation between reef-forming and non-reef-forming species could imply a divergence in the environmental niche, which is likely independent from the evolutionary divergence of "Complex" and "Robust" clades. The diversity of niche relationships observed between the studied scleractinian species, with no pattern related with evolutionary relationships, reflects the complex and poorly understood evolution of Scleractinia.

L. pertusa, M. oculata, S. variabilis and *E. rostrata* are widely considered the main reef-forming species in the central and south Brazilian areas (Kitahara, 2009; Cavalcanti et al., 2017). In this study, we observed that *E. rostrata* had a predicted distribution that overlapped

with *M. oculata* and *S. variabilis* in the South and Central area of Brazil, with all having significantly similar niche. This corroborates their role of reef-forming species in Brazilian waters. It is important to highlight that *E. rostrata*, as well as *S. variabilis*, had a restricted environmental niche (due to inhabit a more restricted range of temperature, depth, POC, AST and regional flux). Thus, *E. rostrata* and *S. variabilis* niches were quite different to the other species niches, showing a non-similar niche with *L. pertusa, Deltocyathus* spp. and *C. debilis*, and with *M. oculata*, respectively. Despite their restricted niche at regional scale, both species are known to be particularly abundant in Campos basin (Cavalcanti et al., 2017). *E. rostrata* and *S. variabilis* are more abundant in the South-west Pacific Ocean, and are infrequently observed in the North Atlantic (Davies and Guinotte, 2011; Roberts et al., 2006).

As discussed for Scleractinia and Octocoralia, substrate habitat type may determine the real distribution, inside the predicted potential distribution areas of the scleractinian species. All reef-forming species together with the non-reef-forming *C. debilis* inhabit hard-bottom substrate, whereas *Deltocyathus* spp. inhabit soft-bottom substrate. For instance, *D. italicus* and *Deltocyathus* sp., solitary species with unattached bases, were registered in both mid and lower slope in Campos basin (Arantes et al., 2009). These cup corals can also create large dense fields, as observed in La Foneca canyon by *Desmophyllum diantus* (Ayma et al., 2019; Lastras et al., 2019), providing hard substrata and increasing local biodiversity. This indicates that diverse kind of VME are likely distributed along the Brazilian continental margin. Particularly in the central and south areas, were a high number of species share habitat suitability (Fig. 2 b - ii and iv).

Species niche properties as niche breadth are indicators of species sensitivity (Kotiaho et al., 2005; Thuiller et al., 2005) and generally, species with a restricted or smaller niche, have been shown to be more sensitive to the loss of habitat caused by disturbances such as those produced by climate change (e.g., fish: Munday, 2004; birds: Seaone and Carrascal, 2008). Given that E. rostrata and S. variabilis had smaller niches than other species in this study, they may be the most sensitive species to variations in environmental conditions. Conversely, those species with a wider environmental niche, such as C. debilis, Deltocyathus spp., L. pertusa and M. oculata may be less sensitive to environmental changes. Despite this assumption, some species could be overly susceptible to changes in one particular environmental factor. For instance, C. debilis is likely to be more sensitive to changes in the dissolved oxygen and silicate concentration than to other factors studied (Fig. 4 d). Specific biological traits, such as acclimatization or adaptive capacity will also influence the species response to environmental changes. For example, L. pertusa has a higher acclimatization capacity to lower temperature than M. oculata (Naumann et al., 2014). However, M. oculata fossils indicate a wider tolerance to changes in environmental conditions than L. pertusa in the Gulf of Cádiz (Wienberg et al., 2009). Specific experimental research of organisms inhabiting different regions are fundamental to understanding their ecological niche, physiology and additional sources of uncertainty that might influence their survival under future environmental disturbances.

4.1. Approach limitations

Whilst habitat suitability modeling has been widely used to determine the potential distribution of deep-sea species (e.g., Davies and Guinotte, 2011; Tracey et al., 2011; Yesson et al., 2012), there are still limitations that should be considered during the modeling approach (Vierod et al., 2014; Anderson et al., 2016). For example, a regularly spaced sampling regime that covers the entirety of the environmental conditions observed in the region of interest is important (Hirzel and Guisan, 2002). Independent species presence and/or absence data is also an important aid for model validation (Anderson et al., 2016). However, deep-sea surveys with this characteristic are extremely limited due to high cost and significant logistical restrictions such as access to both ship-time and high-quality sampling equipment. These limitations lead to a disproportionate effect on sampling quality in some regions, particularly in areas such as the South Atlantic Ocean, where many countries have restricted access to ocean-going research vessels. In this study, the focus on the collection of new region-specific occurrences that were not present within international datasets led to substantial improvements in the quality and utility of predictions in this region. Nevertheless, future surveys must be carried out in order to validate the present predictions, principally in the North and Northeast areas for scleractinian and in the South area to octocorals.

It is essential to note that this study only presents suitable areas, which are statistically likely to contain species presences based upon the environmental data used in the models. There will be other variables or factors that were not included in the analysis which could influence these predictions, such as substrate availability and type. Hard substrate presence is highly variable over small spatial scales and is a strong constraining variable that limits the distribution of many coral species (Davies and Guinotte, 2011; Tracey et al., 2011; Guinotte and Davies, 2014; Mackay et al., 2014). For example, suitable predicted areas for reef-forming species in the flat tops of some seamounts in the South Pacific, were dominated by sand, an unsuitable substrate for most stony coral species settlement (Anderson et al., 2016). Furthermore, the SRTM30 bathymetry data have shown to overestimate suitable area for some deep-sea corals (Marshall, 2011; Ross et al., 2015), particularly in regions were local bathymetric data does not exist (Anderson et al., 2016). Better local surveys and the collection of high-resolution bathymetric data will improve bathymetric representation. This may provide a representation of sea-bed physiographical features of importance for the presence of corals and provide the potential for the elucidation of substrate type, which will substantially improve regional and local habitat suitability models (e.g., Howell et al., 2011; Rengstorf et al., 2013, 2014; García-Alegre et al., 2014; Georgian et al., 2014).

5. Conclusions

The information presented in this study represents the first regional scale habitat suitability modelling effort for CWCs along the Brazilian margin. This study represents a fundamental step in better understanding the distribution of Brazilian CWCs and provides essential information to guide future surveys and conservations plans in the region. For instance, future surveys must be oriented to describe the presence of both, octocorals and scleractinians corals, to validate the predictions, mainly where suitability is observed with no presence records. Based upon the areas of predicted suitable habitats for CWC, we highlight the importance of management plans that combine the oil and gas exploration areas with conservation and mitigation of potential impacts to these communities in Campos Basin, Santos Basin, Vitoria-Trindade chain, Rio Grande rise and Pelotas Basin (South area). Particularly, the Rio Grande Rise area must be characterized in terms of biodiversity composition, since it contains cobalt crusts and a possible future extraction of minerals could impact coral communities that may be currently unknown. The broader habitat suitability distribution of Octocorallia compared to Scleractinia is related to a broader environmental niche breadth, likely due to the variety of physiological adaptations of this group. Non-reef-forming scleractinian species were less restricted by environmental conditions, when compared to reef-forming species, allowing then to potentially inhabit a broader area of the Brazilian continental margin. Most reef-forming species presented similar but not equivalent ecological niche, indicating that management planning, conservation efforts and cruise planning should consider the species individual environmental requirements to be more efficient. For instance, priority consideration may need to be given to S. variabilis and E. rostrata since they had smaller niche breadth and may be more sensitive to changes in the environment.

Declaration of competing interest

The authors declare that there is no conflict of interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.dsr.2019.103147.

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