

Spatial and temporal representation of marine fish occurrences available online

³ Vanessa Pizarro^a, Andrea G. Castillo^{a,b}, Andrea Piñones^{c,d,e,f}, Horacio Samaniego^{a,g,*}

⁶ ^a*Laboratorio de Ecoinformática, Instituto de Conservación, Biodiversidad y Territorio, Universidad Austral de Chile, Valdivia, Chile*

^b*Programa de Doctorado en Ciencias mención Ecología y Evolución, Escuela de Graduados, Facultad de Ciencias, Universidad Austral de Chile, Valdivia, Chile*

⁹ ^c*Instituto de Ciencias Marinas y Limnológicas, Facultad de Ciencias, Universidad Austral de Chile, Valdivia, Chile*

¹² ^d*Centro FONDAP de Investigación en Dinámica de Ecosistemas Marinos de Altas Latitudes (IDEAL), Valdivia, Chile*

¹⁵ ^e*Centro de Investigación Oceanográfica COPAS-COASTAL, Universidad de Concepción, Chile*

¹⁸ ^f*Millenium Institute Biodiversity of Antarctic and Subantarctic Ecosystems - BASE, Chile*

²¹ ^g*Instituto de Sistemas Complejos de Valparaíso, Subida Artillería 470, Valparaíso, Chile*

¹⁸ Abstract

Despite the 243,000 species of marine species described by 2022, our knowledge about the biodiversity in the oceans is still incomplete. This may have dreadful and detrimental effect for the conservation of marine ecosystems under the current anthropization of our biota and the fast pacing climate and global change scenario.

However, a large number of online repositories cataloging, storing and distributing biodiversity information hosting taxonomic information and species occurrence data have emerged recently. FishBase, the Global Biodiversity Information Facility (GBIF) and the Ocean Biodiversity Information System (OBIS) are part of these publicly available repositories representing a variety of sources that have exploded in number. However, despite the incredible accumulation of biodiversity records, not all the information is really useful, nor does it represent any new knowledge regarding global species richness patterns.

In this study we evaluated the spatial and temporal representativeness of the records of marine fishes (order Actinopterygii) available in the GBIF

²⁴*Corresponding author

²⁷Email address: horacio@ecoinformatica.cl (Horacio Samaniego)

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and OBIS global repositories. We provide a methodological framework based on a set of non-parametric estimators to calculate species richness from incidence data, using hexagonal grids as sampling units overlapped on the marine bioregions worldwide.

Using standard ecological and spatial analysis tools, we identify regions that are adequately represented in terms of available records and therefore have more reliable data, as well as regions with few records that do not represent current species richness. We overlap these results with the location of marine protected areas and fishing exploitation zones to understand the anthropogenic effect on marine ichthyofauna. We additionally evaluate hypotheses regarding the taxonomic, geographic, and temporal distribution of information biases to deepen our current understanding of public records of species occurrences worldwide.

Considering that more than 40 years of information was analyzed, the results showed that on a global scale, the primary data on marine fish available on GBIF and OBIS platforms are still far from being representative and complete. Only 1.14% of the records were useful for our analyses. In addition, we found that the information seems to be biased towards coastal areas, regions close to developed countries and areas where there is a large fishing activity. Finally, the best represented species and families are those with a small body size, which use shallow habitats and have commercial cultural value.

Keywords: Ecoinformatics, Ecological Information Biases, Marine Fish, Spatial and Temporal Representativeness, Species Richness

1. Introduction

Currently, the more than 243,000 species included in the World Register of Marine Species database ([WORMS, 2022](#)) suggests that only 11 to 78% of all marine species have been discovered revealing a striking picture of vastly incomplete knowledge that may have serious implications for marine conservation ([Luypaert et al., 2020](#)). Moreover, ongoing climate change represents one of the greatest threats to biodiversity ([Malhi et al., 2020; Turner et al., 2020](#)) and has already been documented to modify the distribution of marine species ([Lenoir et al., 2020](#)). Some of the described effects considers the

invasion of non-native species leading to massive species turnover that may lead to the local extinction of large proportions of species (Cheung et al.,
3 2009).

While species richness is often used to represents diversity patterns, species richness is, in itself, an aggregate variable subsuming the variety of life (Marquet et al., 2004). Hence, several attempts have focused on the development of more encompassing indices sparking interesting scientific debates to describe ecological heterogeneity (Tuomisto, 2011; Moreno and Rodríguez,
6 2011; Daly et al., 2018). However, scientific literature seem to have opted to shift focus to the consequences of biodiversity loss by fostering the usage of new terminology to provide hands-on concepts such as species inventory,
9 taxonomic inventory or inventory completeness designed to convey sharper messages to policy makers summarizing the richness of biodiversity (Pereira et al., 2013; Butchart et al., 2010). Still, while scientists have debated the use
12 of biodiversity terminology, species richness provides a succinct, and easy-to-handle description of the variability across several other quantities describing the biota in space and time (Appeltans et al., 2012), and is an essential feature
15 to understand how diversity changes under the impact of natural and anthropogenic factors on biomes, regions and ecosystems (Troia and McManamay,
18 2017; Magurran and McGill, 2011).

Likewise, biodiversity can also be assessed through life history traits, which are modulated by both evolutionary factors and the variation in habitats and ecosystems (Neigel, 1997; Hutchings and Baum, 2005). We now
21 know that biodiversity is more likely an expression of the heterogeneity of such life history traits. Alò et al. 2021, for example, shows that while some of the fish diversity is certainly due to environmental processes, a large fraction
24 of such richness variance is also determined by evolved life history traits related, for example, to migratory habits. Therefore, evaluating how life history traits impact richness metrics should deepen our understanding of fish
27 diversity patterns.
30

While still short of having a robust and standardized biodiversity infrastructure ([Heberling et al., 2021](#)), a great diversity of online repositories with taxonomic information and species occurrences data exist. Among the most important databases hosting marine information are FishBase, a platform that hosts information on the taxonomy of fish, their ecology, trophic information, habitat and history of uses dating back to more than 250 years ([Froese and Pauly, 2000](#)); the Global Biodiversity Information Facility (GBIF), a platform that stores and allows for the free access to species occurrence records from around the world. GBIF is currently one of the repositories hosting the largest amount of such data in the world ([Telenius, 2011; GBIF: The Global Biodiversity Information Facility , 2021](#)); and finally Ocean Biodiversity Information System (OBIS), which houses data on the occurrence and abundance of species from exclusively marine environments ([OBIS: Ocean Biodiversity Information System, 2021](#)). Records entered in these repositories are often used for research related to biodiversity assessment, taxonomic reviews, red listing of threatened species, species distribution, and generation of ecological niche models, among others ([Yesson et al., 2007](#)). GBIF currently offers more than 1.62 billion occurrence records and OBIS more than 63 million, which increase considerably each year ([GBIF: The Global Biodiversity Information Facility , 2021; OBIS: Ocean Biodiversity Information System, 2021](#)).

The records of both platforms come from a wide variety of sources collected following different methodologies at different temporal and spatial scales introducing a great variety biases ([Beck et al., 2014; Zizka et al., 2020](#)). Among these, three main types of biases have been described: (i) taxonomic, this occurs when some species and/or families are better sampled than other rarer species ([Chandler et al., 2017](#)); (ii) geographic, when data input is unevenly distributed across geographic regions and may prove to obscure inter-region comparisons ([Yang et al., 2013; Yesson et al., 2007](#)); and (iii) temporal, which may be prevalent when comparing different time periods as

data coverage is unevenly distributed over time (Chandler et al., 2017; Yang et al., 2013). While these biases introduce some uncertainty regarding reliability of species richness descriptions obtained from online platforms (Beck et al., 2014; García-Roselló et al., 2015), they have largely been used to provide an extensive overview of macro-ecological patterns of distribution not available otherwise (Mora et al., 2008; Troia and McManamay, 2017).

Still, identifying how sampling effort is distributed across space and time is a necessary step to interpret biodiversity patterns and reduce biases as understanding the distribution of our biota is essential to design protection efforts. This may be achieved through different weighting schemes for records in areas with sufficient sampling that provide a more reliable contribution compared to underrepresented regions (Phillips et al., 2009; Hortal et al., 2008; Yang et al., 2013).

We here assessed the spatial and temporal representativeness of marine fish records available in the global GBIF and OBIS repositories at the level of marine bioregions in order to pinpoint the location of records that best quantify the diversity of marine fishes. The result is a spatial representativeness analysis that we then overlay on marine conservation areas (UNEP-WCMC and IUCN, 2022) and fisheries exploitation areas (FAO, 2014) to learn whether marine conservation efforts, as well as large fisheries, are located in areas of high species richness or in areas of insufficient data coverage.

Finally, we also analyzed the potential effect that some attributes could have on the incidence of more records in global database repositories. Specifically, we evaluated three hypotheses related to body size, habitat depth and commercial use. The underlying hypotheses are that a better representation in the online platforms may be due to the over sampling of larger fish, caused by its easy identification; that shallow areas provide easy access to sampling; and economic and commercial interest have elicit a larger representation of culturally relevant species among online biodiversity repositories.

2. Methods

2.1. Species data

We use all recorded occurrences from the order Actinopterygii hosted in GBIF and OBIS repositories (GBIF.org, 2021; OBIS.org, 2021). Following Alò et al. (2021), evolutionarily older taxa, such as Cephalaspidomorphi, were excluded from this analysis. The libraries *rgbif* and *robis* of the statistical package R were used for data extraction (Chamberlain, 2017; Provoost and Bosch, 2020; R Core Team, 2018). To minimize errors associated with the public usage of GBIF and OBIS repositories, we curated the dataset following Zizka et al. (2020) and filtered the dataset by the columns labeled “scientific name”, “family”, “year”, “Longitude” and “Latitude”. We retained all taxonomic information down to the species level. Any record with NA values was removed. We also removed any duplicated record with identical latitude and longitude as well as any record collected before 1980 (see Alò et al., 2021; García-Roselló et al., 2015). Each record was further assigned to a marine bioregions following Costello et al. (2017). Spatial data wrangling and plotting was performed with the aid of the following libraries: *sf*, *dplyr*, and *cartography* (Giraud and Lambert, 2016; Pebesma, 2018; Wickham et al., 2021). We finally labeled and removed any exotic species record using the *distribution()* function provided by the *rfishbase* library (Boettiger et al., 2012; Froese and Pauly, 2021). To limit our analysis to species occurring within their native range each record was checked against the classification of FAO fisheries area for consistency (FAO, 2014).

2.2. Data Analysis by Bioregion

Once the database was cleaned, a subset of the data was created for each of the 30 bioregions. For each bioregion, a count of records, species and families was made, and the Shannon diversity index was calculated using the *vegan* library in R (Oksanen et al., 2020).

2.2.1. Spatial Representativeness Analysis

To assess the spatial representativeness of the data, bioregions were gridded into hexagonal cells of equal area to maximize fit to bioregions areas using a cylindrical equal area projection and the World Geodetic System of 1984 Alò et al. (2021). These $1^\circ \times 1^\circ$ hexagonal lattice yielded 57,067 cells in total. We evaluated, in the appendix, two additional spatial resolutions, of 5° and 10° lattice with a total of 3,029 and 953 cells respectively, in order to analyze different biodiversity macropatterns (Tittensor et al., 2010). The expected species richness (S_{exp}) was computed as the average between three non-parametric richness estimators so that $S_{exp} = \frac{1}{3} \sum_i^3 S_i$, where S_i is Chao2 (S_{chao}), Bootstrap ($S_{bootstrap}$) and Jackknife 1 ($S_{jackknife1}$) (see Magurran and McGill, 2011, for individual definition of indices). Such averaging seeks to minimize biases and potential errors of under- or over-estimation by using a single richness estimator following the work of (Mora et al., 2008; Troia and McManamay, 2017).

We then produced a spatial representativeness index (SRI) by comparing the observed richness (S_{obs}) per cell to S_{exp} (Troia and McManamay, 2017), $SRI_i = \frac{S_{obs}}{S_{exp}}$. This index indicates the degree of representativeness of records to quantify the actual species richness in each cell (i). Its value ranges from 0 to 1, where 0 represents an unsampled cell and 1 represents a fully sampled one.

Because SRI somehow shows how databases depict the actual species richness, we may further classify SRI into four classes labeled by the levels of species richness knowledge they represent. Some cell may show very *few* knowledge with $SRI \in (0, 0.60)$. Conversely, others may show to have a *sufficient* species diversity knowledge level for a complete representation of species diversity if $SRI \in (0.60, 0.85)$. While others, in turn, will have an *adequate* representativeness level if $SRI \in (0.85, 1.00)$. Cells with one or *no records* were also considered as an independent class, as well as cells with a single record in order to identify those cells with insufficient records for SRI

estimation.

2.2.2. Temporal Representativeness Analysis

We generated plots of cumulative records over time to analyze the temporal distribution of data records for each bioregions. Accumulation curves for the 30 bioregions were calculated based on the observation records and the year of collection. We assess the completeness of the sample by evaluating the final 10% tip of the curve using a linear fit after rescaling S to create statistically comparable slope units. Slopes close to 0 indicate sufficiently sampled bioregions, while slopes closer to one are indicative of insufficient sampling efforts across time.

2.2.3. Gap Analysis

We overlaid the spatial representativeness map (§2.2.1) with shapefiles of Marine Protected Areas (MPA) (UNEP-WCMC and IUCN, 2022) and fishing exploitation areas reported by (FAO, 2014). The superposition of these layers allowed us to calculate the extent of protection offered by MPA for each bioregions on a cell basis, and the extent of cells in designated fishing zones. This exercise allows to jointly assess the relationship between two opposing human impacts and current uncertainties about marine fish diversity.

2.2.4. Bias Assessment

The evaluation of potential biases generated by body size, habitat depth and cultural value of species (§2.1) was assessed from the fishbase database (Froese and Pauly, 2021). Size frequencies were determined using 80 cm intervals and ranges of habitat depth were determined according to the classification of oceanic layers (i.e. epipelagic = 0 - 200 m, mesopelagic = 200 - 1,000 m, and bathypelagic= 1,000 - 4,000) (Costello et al., 2010). Parametric correlation analysis was employed describe the relationship between the frequency of representation, using a logarithmic transformation, and the

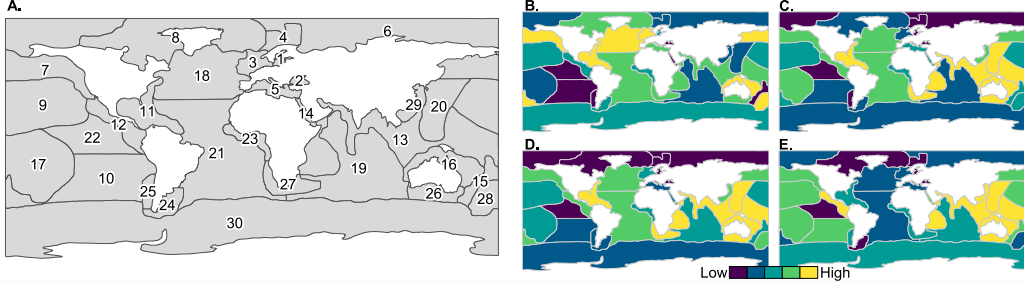


Fig. 1: Marine bioregions and spatial diversity distribution used in this study. **A.** The 30 marine bioregions from Costello et al. (2017) used in this study. Number are identification labels in Table 1. **B.** Records by bioregion; **C.** Overall species richness across bioregions; **D.** Family richness; and **E.** Shannon diversity index. Note that values in **C-E** have been normalized for display purposes. See Table 1 for actual values.

body size and habitat depth, while a simple pie chart shows the frequency of cultural values associated to the data.

- 3 All data and scripts are available (see Appendix A).

3. Results

3.1. Records by Bioregions

6 Approximately 1.14% of the total published occurrences in the order
Actinopterygii were retained in our analysis. That is, from the 71,670,596
downloaded records off the GBIF and OBIS repositories, 820,004 were con-
9 sidered useful (see Appendix A). This subset consisted of 10,371 species in
361 families. The most represented families in our dataset are Scombridae,
Pleuronectidae and Gadidae with 103,762, 57,018 and 52,079 records
12 respectively. The species with the largest representation frequency are *Hip-*
poglossoides platessoides, *Mola mola* and *Coryphaena hippurus* with 30,885,
21,042 and 21,089 records.

15 The analysis at bioregion level (Table 1) shows a large variability. The
count of records varies across three orders of magnitudes, that is from $2.68 \times$
 10^5 records in the Caribbean Sea and Gulf of Mexico (11) down to $1.02 \times$
18 10^2 in the Black Sea (2). The bioregion with the largest species richness

Table 1: Area (1,000 km²) and counts of records, species richness, family richness and Shannon diversity for each bioregion. The largest values for each column is highlighted.

ID	Bioregion	Area	Records	Species	Families	Shannon
1	Inner Baltic Sea	415	8,902	72	30	2.46
2	Black Sea	537	102	37	22	3.21
3	NE Atlantic	2,053	87,377	310	104	3.90
4	Norwegian Sea	1,132	3,046	93	35	2.16
5	Mediterranean	2,859	12,532	372	101	3.39
6	Arctic Seas	10,276	2,506	114	23	3.90
7	North Pacific	12,974	78,070	839	156	4.50
8	North American boreal	8,001	9,709	162	48	2.99
9	Mid-tropical N Pacific Ocean	32,685	9,310	615	127	4.59
10	South-east Pacific	21,952	386	190	89	4.97
11	Caribbean and Gulf of Mexico	8,427	268,066	1,703	209	4.49
12	Gulf of California	6,184	7,639	885	148	5.93
13	Indo-Pacific seas and Indian Ocean	37,090	16,967	2,947	215	6.93
14	Gulfs of Aqaba, Aden, Suez, Red Sea	830	926	352	72	5.51
15	Tasman Sea	3,592	1,003	380	120	5.36
16	Coral Sea	7,658	40,107	2,929	249	6.75
17	Mid South Tropical Pacific	23,418	6,083	811	123	5.18
18	Offshore and NW North Atlantic	16,012	130,994	897	190	3.46
19	Offshore Indian Ocean	31,076	1,263	337	116	4.06
20	Offshore W Pacific	10,291	6,363	1,839	232	6.81
21	Offshore S Atlantic	41,435	11,960	990	188	3.79
22	Offshore mid-E Pacific	13,815	687	79	37	3.04
23	Gulf of Guinea	3,325	6,816	384	138	3.95
24	Argentina	2,665	8,701	115	52	2.83
25	Chile	1,739	250	100	54	4.36
26	Southern Australia	3,824	15,643	1,011	201	5.75
27	Southern Africa	4,371	19,954	1,142	210	4.16
28	New Zealand	6,293	53,879	558	154	3.66
29	North West Pacific	2,457	1,767	869	182	6.46
30	Southern Ocean	62,161	8,996	294	57	3.98

and diversity index is the the Indo-Pacific Seas and Indian Ocean (13) with 2.95×10^3 recorded species and a Shannon index of 6.93 followed by the
3 Coral Sea bioregion (16) with 2.93×10^3 species and a Shannon index of 6.75. Likewise, the Coral Sea also presents the largest number of families. It
6 is interesting to note that, while being the largest bioregion (i.e. in km²), the
Southern Ocean show the fewest number of records and the lowest number
9 of species and families across all bioregions. Black Sea (2) and Norwegian
respectively. Fig. 1 illustrates the location of the 30 marine bioregions and
their respective richness and diversity values.

3.2. Geographic Analysis

12 Fig. 2 shows the cell classification according to SRI (§2.2.1). As expected,
no bioregion is completely sampled at the 1° scale resolution. In fact, at this
resolution scales, large empty regions with no records are observed. The
15 bioregions with the largest area classified as *Adequate* are the Northeast Atlantic (3) (37.53%), the Caribbean and Gulf of Mexico (11) (29.26%) and the
Inland Baltic Sea (1) (24.37%). It should be noted that such cells are mostly
18 correspond from coastal areas in the northern hemisphere. On the other
hand, the bioregions that present a greater surface without records corre-
spond to the Southeast Pacific (10) (96.3%), the Arctic Sea (6) (94.9%), and
21 the Southern Ocean (30) (93.7%). While the bioregions with the larger sur-
face with sufficient records are the Gulf of Guinea (23) (32%), the Norwegian
Sea (4) (22.3%), and the Gulf of California (12) (21.6%). Additional results
24 for 5° × 5° and 10° × 10° spatial resolution grids are available in [Appendix B](#).

3.3. Temporal Analysis

27 Bioregions show similar trends of data accumulation across the four decades
analyzed here (Fig. 3). While a significant increase is apparent in the time
period between 2005 and 2010, such increase is not significant for 14 out the

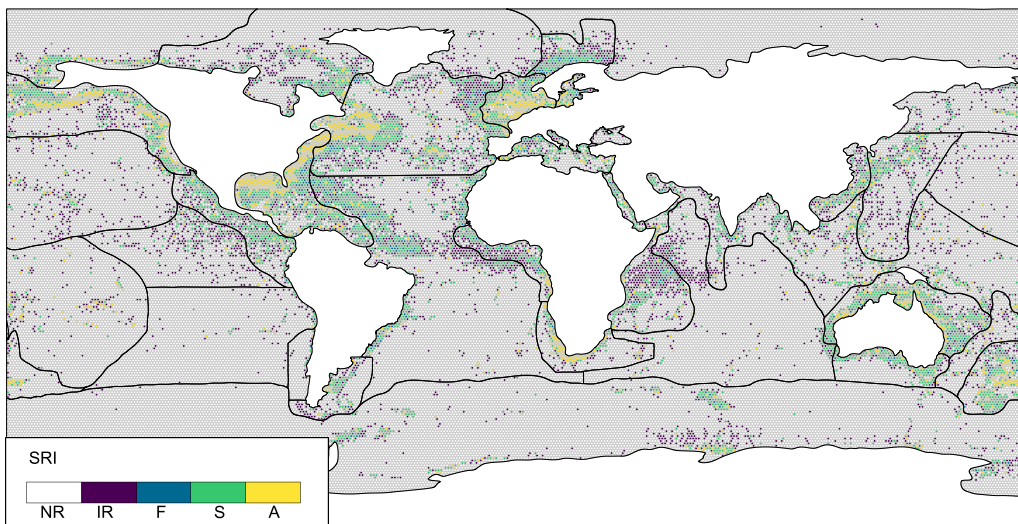


Fig. 2: Spatial Representativeness Index (SRI) in 1° hexagonal lattice. Values in the table below indicates the surface area as a percentage of each bioregion for every SRI category (see §2.2.1). **ID** is the identification number given to each bioregion (Table 1). **A** are cells with an adequate representativeness of species richness (i.e. $SRI > 0.85$). **S** are cells considered as having a sufficient representativeness (i.e. $SRI \in (0.60, 0.85)$). **F** cells are cells with few records and are thus not considered to be representative of actual species richness (i.e. $SRI \in (0, 0.6)$). **NR** as cells with no records ($SRI = NA$).

30 bioregions. The Caribbean and Gulf of Mexico (11) is the bioregion with the largest increases in data contribution to the dataset, while the Black Sea

- ³ (2) is the bioregion with the lowest rate of data contribution in the 40 years span between 1980 and 2020. (See Appendix C for further analysis).

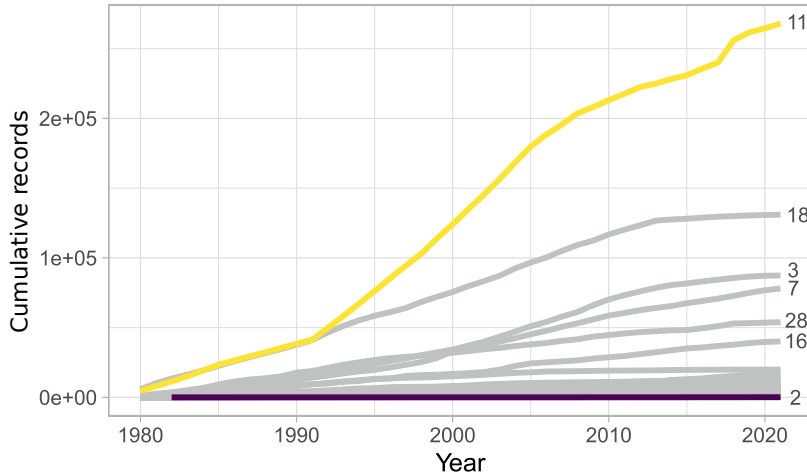


Fig. 3: Records accumulation rate for each bioregion across the four decades analyzed. The blue line is the accumulation of fish records in the Caribbean and Gulf of Mexico bioregion (11) and the red line shows the accumulation rate in the Black Sea (2). Numbers as the end of each timeseries correspond to the bioregion ID in Table 1.

We categorize the slopes of the final 10% of each accumulation curves in

- ⁶ Fig. 4. Fourteen bioregions show a slope less than 1. The Mediterranean Sea (5) stands out with the lowest slope value (0.47), while the Black Sea (2) is the bioregion with the steepest final slope (3.13).

⁹ 3.4. Gap analysis and fishing exploitation areas

The bioregions with the largest area covered by protected areas are the Coral Sea (16), the northeast Atlantic (3) and New Zealand (28) covering a ¹² 37.3, 17.4 and 16% of their respective areas. Regarding the sampling level of these bioregions, the Offshore Indian Ocean (19); Gulf of Aqaba, Aden, Suez, Red Sea (14); and Coral Sea (16) are the bioregions with the highest ¹⁵ percentages of cell sampled as *Adequas* inside of their protected areas (83,

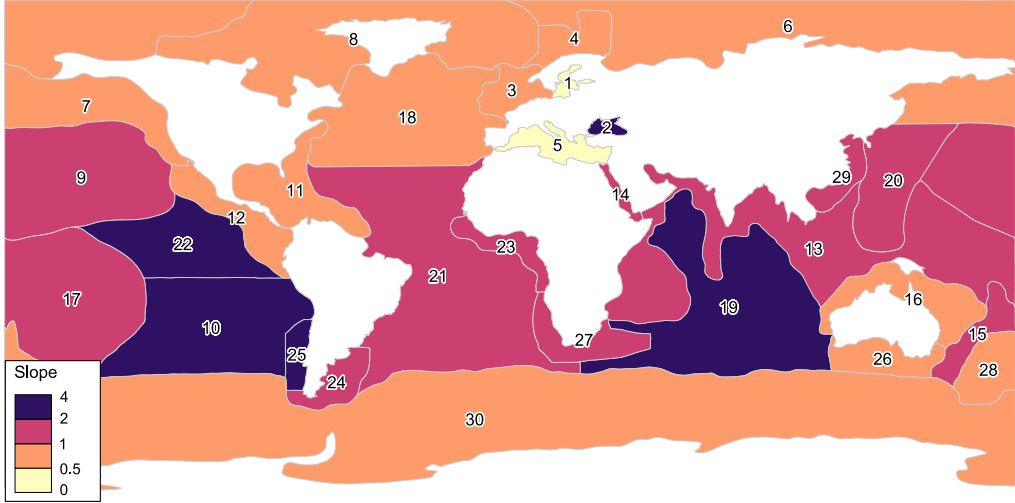


Fig. 4: Graphical representation of the slope values of the species accumulation curve for each bioregion. The slope corresponds to the final 10% of the species accumulation curve. See §2.2.2 for details regarding the analysis.

63.8 and 59.8% respectively). While the Arctic Seas (6), North American boreal (8) and Mid South Tropical Pacific are the bioregions with protected areas with the highest percentage of cell with no records (86.2%, 83.8% and 81.2%). (See Appendix D).

FAO areas with the largest area categorized as *Adequate* correspond to the northwest Atlantic (22.1%), Northeastern part of the Pacific Ocean (14.6 %), and Western part of the Atlantic Ocean (12.6 %) (Table 3). These FAO areas correspond to regions of the Pacific Ocean (North Pacific, North West Pacific, Mid-tropical N Pacific Ocean and Indo-Pacific seas and Indian Ocean, as well the Gulf of California and Caribbean and Gulf of Mexico). Largest FAO areas with *NR* correspond to the Antarctic part of the Pacific Ocean, the Antarctic part of Atlantic Ocean and Southeastern part of the Atlantic Ocean in the Southern Ocean, Offshore S Atlantic and Southern Africa.

Table 2: Results of overlapping Marine Protected Areas and SRI grid. ID is the identification number given to each bioregion (see Table 1 for bioregion names). Area corresponds to the percentage of surface area covered by marine protected areas. NR is the percentage of cells with *No Records*; IR is the percentage of cells with *Insufficient records*; F is the percentage of classified cells with *Few* records; S, the percentage of classified cells with *Sufficient* records, and A, the percentage of classified cells with *Adequate* records. The highest values for each column is highlighted.

ID	Area	NR	IR	F	S	A
1	0.03	2.38	4.30	5.22	49.08	39.01
2	12.89	26.71	27.61	10.49	35.19	0.00
3	9.74	3.23	1.35	0.94	40.86	53.62
4	0.15	11.16	19.63	5.18	56.69	7.34
5	0.09	5.71	6.77	11.20	47.95	28.37
6	5.01	86.16	5.97	0.02	4.65	3.20
7	0.00	26.48	4.24	1.26	23.77	44.25
8	1.23	83.82	7.77	0.62	6.70	1.11
9	0.69	69.58	15.77	0.00	6.40	8.25
10	17.36	73.51	24.58	0.00	0.80	1.11
11	0.28	20.13	6.25	3.44	29.98	40.21
12	0.83	0.33	1.23	8.85	61.35	28.25
13	0.45	50.52	11.94	0.88	25.17	11.50
14	0.25	8.87	0.00	1.59	25.65	63.88
15	4.06	57.18	15.27	0.00	7.29	20.26
16	16.00	3.10	0.49	1.10	35.52	59.79
17	0.20	81.19	11.43	0.00	3.07	4.31
18	4.91	35.87	16.63	0.77	25.64	21.09
19	2.78	11.88	0.74	0.00	4.34	83.04
20	2.56	34.06	9.68	6.85	35.31	14.10
21	3.79	51.06	19.35	0.38	21.35	7.86
22	0.16	41.98	27.54	0.00	23.22	7.26
23	13.83	9.92	4.92	7.98	61.33	15.85
24	0.21	40.86	20.64	0.24	23.97	14.29
25	0.07	65.27	0.50	0.05	1.29	32.89
26	0.28	30.04	12.83	6.89	38.62	11.63
27	1.45	16.78	5.75	0.15	18.71	58.62
28	0.00	45.36	2.25	0.00	13.71	38.67
29	37.29	20.49	17.05	2.68	42.50	17.29
30	1.66	64.05	7.12	4.89	19.86	4.08

Table 3: Results of overlapping FAO fishery exploitation areas and SRI grid. The surface area corresponding to each bioregion, and the percentage of surface area of each classification. Area is in thousands of km²; NR is the percentage of cells with *No Records*; IR is the percentage of cell with *Insufficient Record*; F is the percentage of classified cells with *Few* records; S, the percentage of classified cells with *Sufficient* records, and A, the percentage of classified cells with *Adequate* records. The highest values for each column is highlighted.

FAO Area Name	Area	NR	IR	F	S	A
Arctic Sea	4,085.78	93.22	3.13	0.29	2.61	0.75
Northwestern part of the Atlantic Ocean	873.55	31.19	11.66	5.69	29.37	22.08
Northeastern part of the Atlantic Ocean	3,222.88	66.29	12.54	2.55	13.63	4.99
Western part of the Atlantic Ocean	1,285.13	30.84	13.09	7.91	35.60	12.55
Eastern Central part of the Atlantic Ocean	1,207.71	52.61	24.09	3.44	18.37	1.19
Mediterranean Sea and the Black Sea	308.53	46.39	15.43	5.24	24.77	8.17
Southwestern part of the Atlantic Ocean	1,730.57	82.49	5.85	1.69	8.55	1.42
Southeastern part of the Atlantic Ocean	1,765.33	89.92	4.19	0.15	2.13	3.61
Antarctic part of the Atlantic Ocean	2,310.48	93.31	2.80	0.20	2.93	0.76
Western part of the Indian Ocean	2,620.90	72.45	16.11	1.03	8.51	1.89
Eastern part of the Indian Ocean	3,028.72	85.40	4.69	0.82	7.39	1.70
Antarctic and Southern of the Indian Ocean	1,977.29	85.71	7.76	0.56	4.33	1.64
Northwestern part of the Pacific Ocean	2,259.45	73.55	12.40	0.94	10.32	2.79
Northeastern part of the Pacific Ocean	967.54	55.13	12.65	1.34	16.26	14.62
Western Central part of the Pacific Ocean	2,963.11	70.45	12.56	0.43	11.58	4.98
Eastern Central part of the Pacific Ocean	4,140.60	79.36	11.30	0.31	6.94	2.09
Southwestern part of the Pacific Ocean	3,096.68	85.04	4.42	0.97	6.40	3.17
Southeastern part of the Pacific Ocean	2,997.11	91.16	6.00	0.10	2.30	0.44
Antarctic part of the Pacific Ocean	2,361.33	93.47	4.57	0.21	1.42	0.33

3.5. Evaluation of Biases

We evaluated biases for body size, habitat depth, and cultural value for 3 10,371 marine fish species identified in our database (§3.1).

3.5.1. Body size

The range 0-80 cm is the most frequently occurring size length. Three species stand out with the highest numbers of records, *Scomber scombrus*, *Lagodon rhomboides* and *Mallotus villosus* with 20,995, 19,563 and 13,609 records respectively. These species are distributed mainly in the Northeast Atlantic (3) and Offshore and Northwest North Atlantic (18) bioregions. While the families that accumulate the greatest number of records correspond to Sparidae, Scombridae and Labridae with 24,837, 21,719 and 21,035

records. These families are mainly distributed in the Caribbean Sea and Gulf of Mexico and the Northeast Atlantic.

3 3.5.2. Habitat depth

The most frequent depth range is between 0-838 m (i.e. epipelagic and mesopelagic zones), and the species with the highest number of records are *Mola mola*, *Coryphaena hippurus* and *Lagodon rhomboides* with 21,089, 21,042 and 19,563 occurrences in the databases. These species are distributed mainly around the Caribbean Sea and Gulf of Mexico (11) bioregions, as well as the following bioregions: Offshore and NW North Atlantic (18) and the South Atlantic Coast (21). The families that accumulate a greater number of records correspond to Scombridae, Gadidae, Sparidae with 63,572, 38,876 and 30,041 records. These are mostly distributed in the northern hemisphere. That is, the Caribbean and the Gulf of Mexico (11), Offshore and NW of the North Atlantic (18) and part of the South Atlantic Ocean Coast (21) bioregions.

Figure 5 shows that body size and habitat depth have a negative correlation with the frequency of records.

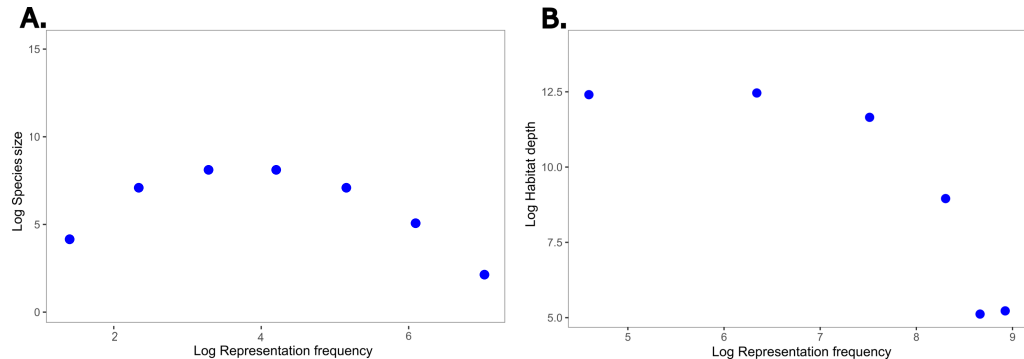


Fig. 5: **A.** Relationship between marine fish representation frequency ($\log_{10} a$) in GBIF and OBIS and habitat depth. **B.** Relationship between the frequency of representation of marine fishes ($\log_{10} a$) in GBIF and OBIS and the size of the species. The dotted line is only there to highlight the negative trend between variables.

3.5.3. Cultural value

Finally, when analyzing the most frequent cultural value represented across our dataset (Fig. 6), “Commercial” use of the species emerges as the most important with a 73.4% among records, followed by the category “No interest” (5.03%), and “Subsistence fishing” (3.08%).

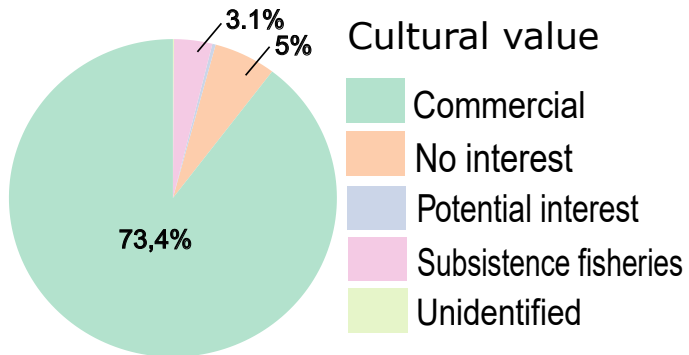


Fig. 6: Frequency of marine fish representation in GBIF and OBIS according to importance of cultural use.

6 4. Discussion

Our work provides a methodological framework based on a set of non-parametric estimators to quantify the potential number of species from incidence data (Chao et al., 2009). We used hexagonal grids that fit the geographic reality of marine ecosystems, and we placed special emphasis on cleaning the occurrence data in their taxonomy (Jin and Yang, 2020) and any potential input errors associated with large and massive datasets (Zizka et al., 2020). This led us to focus on only evaluating marine species in the order Actinopterygii (Alò et al., 2021).

Publicly accessible occurrence records are growing rapidly, partly due to the significant advances in ecoinformatics (Lenoir et al., 2020; Oliver et al., 2021). These databases harbor a growing variety of sources, including museum specimens, field observations, acoustic and visual sensors, and citizen

science efforts (Amano et al., 2016). However, despite the incredible accumulation of biodiversity records, not all the data is really useful, nor does 3 it represent new insights into the distribution of species (Bayraktarov et al., 2019; Zizka et al., 2020). That is why a systematic evaluation of the integrity and coverage of this information is required (Troia and McManamay, 2017).

6 There is an extensive bibliography that evaluates the record quality available for different taxonomic groups. Some examples are: legumes on a global scale (Yesson et al., 2007), lepidoptera from Great Britain and woody plants 9 in Panama (Chao et al., 2009), global marine biodiversity (Tittensor et al., 2010), vascular plants in China (Yang et al., 2013), marine fish on a global scale (Mora et al., 2008; García-Roselló et al., 2015), freshwater fish in the 12 USA (Troia and McManamay, 2017; Pelayo-Villamil et al., 2018), and terrestrial mammals on a global scale (Oliver et al., 2021), among many others. Assuming that that not all data available in these repositories are useful 15 for biodiversity analyses, several efforts have proposed parametric and non-parametric estimators for data cleaning and species richness analysis, Mode- 18 stR (García-Roselló et al., 2013), KnowBr (Lobo et al., 2018), and RWizard (Guisande and Lobo, 2019) among these.

Regarding the units of analysis, here we estimate the species richness at the grid level in order to obtain more uniform results on the distribution of 21 occurrences and avoid overestimating the SRI for marine bioregions (Pelayo- 24 Villamil et al., 2018). In addition, we evaluated two additional grid sizes (5°x5° and 10°x10°), and like other studies, our results show that the coarser 27 the resolution used, the greater the overestimation is, in terms of area. That is, the richness index will indicate that a large area is, indeed, well sampled when in reality, the occurrence records could in fact be groups in a very small 30 area. On the contrary, the finer the scale of analysis, the more localized and deficient the sampling is (Tittensor et al., 2010; García-Roselló et al., 2015; Meyer et al., 2015; Troia and McManamay, 2016, 2017). In view of the above, we recommend using grids that really allow observing macro-ecological

patterns, especially in coastal regions, which may be underrepresented when using lower resolution or square grids (Pelayo-Villamil et al., 2018).

3 Considering that more than 40 years of data were analysed, our results demonstrated that on a global scale, the primary marine fish data available on the GBIG and OBIS platforms are still far from being representative
6 and complete. Compared with other studies evaluating the same taxonomic group (Mora et al., 2008; García-Roselló et al., 2015), although we obtained similar macroecological patterns, only 1.14% of the records extracted from
9 both repositories were useful for our analyses. The large percentage of the occurrences presented input errors or did not have the necessary data to generate a reliable analysis (Yesson et al., 2007; García-Roselló et al., 2014).

12 We also found evidence of strong information biases in the records explored. On the one hand, when analyzing the families and species with the greatest representation, they coincide with groups of fish of commercial interest, demonstrating the existence of **taxonomic bias** of the data
15 (Melo-Merino et al., 2020). This is the case of the families Scombridae, Pleuronectidae and Gadidae, which include species of nutritional importance
18 such as tuna, cod, haddock, among others (Cohen et al., 1990). The same is true for the species with the largest number of records, *H. platessoides* (Pleuronectidae), *C. hippurus* (Coryphaenidae), and *M. mola* (Molidae), the
21 first two are species exploited by the fishing industry, with the exception of sunfish (*M. mola*) which has a wide distribution and is mostly associated with scientific and recreational interest (Pope et al., 2010).

24 The unequal contribution of data at the spatial level is another factor that must be considered to work with data available on ecoinformatic platforms. There is a clear preference for certain regions and/or ecosystems as a result of
27 geographical bias. The literature indicates that the highest data contribution rates correspond to developed countries (Yesson et al., 2007; Chandler et al., 2017), and those coastal regions with better road connectivity (Chandler et al., 2017; Melo-Merino et al., 2020). This information uncertainty is also

particularly prevalent in under-sampled marine habitats, such as the deep sea (Webb et al., 2010). Our results coincide with what is described in the literature, regardless of the size of the grid that was used to generate the analysis, the bioregions that include the Atlantic, the Caribbean and the Gulf of Mexico, and the Baltic Sea are the regions with the highest number of area sampled as *Adequate* associated mainly with coastal areas. However, the number of cells with insufficient data to generate a unbiased diversity analysis, is also worrisome. For instance, our results show that these cells are distributed in more internal areas of the bioregions, zones where sampling is likely to be more difficult. While the bioregions that include the South and Southeast Pacific (including the southern coast of South America), the Southern Ocean, and the Arctic Seas are the regions with the least spatial representativeness of records, the proportion of cells without records (*NR*) exceeds 90%. This large area without samples will make any attempt to describe species richness and distribution highly unreliable in these bioegions (Yang et al., 2013; Troia and McManamay, 2017). The marine regions that include the water column, the seabed and the subsoil beyond the limits of jurisdiction of countries cover almost half of the Earth's surface and support a great abundance and diversity of life (Visalli et al., 2020). Even so, when examining the marine ichthyofauna occurrence data, these represent the least sampled areas.

Finally, the time bias of the data is also present in our study. Diametric differences in species identification and sampling methodologies over the decades have resulted in the production of databases of variable quality. However, the current era is characterized by more accurate data thanks to improvements in individual capture and identification tools (Costello et al., 2015; Jin and Yang, 2020). For these reasons our approach considers occurrence records since 1980, however, the coverage of occurrence data is uneven over time when comparing between marine bioregions. Despite evaluating more than four decades of data, still 46% of marine bioregions have insuffi-

cient sampling efforts. Not surprisingly, the Caribbean and Gulf of Mexico (11) bioregion is the region with the largest input of data, demonstrating
3 once again that geographic sampling bias has strong effects on spatial predictions of species richness (Yang et al., 2013). Future sampling efforts should focus on bioregions at low or equatorial latitudes, areas where biogeographic
6 studies show that marine biodiversity is concentrated (Costello et al., 2017).

All the biases that we have described, added to the inherent problems in data capture, foster and deepen various information gaps that affect the effective spatio-temporal quantification of biodiversity (Magurran and McGill, 2011). In this study, we have overlapped our estimates of species richness with the global marine protected areas declared up to the beginning of the
12 year 2022 (UNEP-WCMC and IUCN, 2022), and the areas of fishing exploitation reported by the FAO (FAO, 2014). This exercise demonstrates the importance of public databases that can faithfully reflect the taxonomic
15 and biogeographical knowledge available for each region of the world (Pelayo-Villamil et al., 2018). Our results indicate that North West Pacific bioregion (19) has the largest area covered by marine protected areas. However, its
18 percentage of adequately sampled cells is low compared to other bioregions. This latter result is of certain concern as this bioregion is considered a conservation hotspot among other bioregions such as the Coral Sea (16), a bioregion
21 with a relatively large percentage of adequately sampled cells (Ramírez et al., 2017). However, we found a low proportion of well-sampled cells in both regions, demonstrating the existence of important information gaps, at least
24 for fish of the order Actinopterygii. We emphasize the need to correct these information gaps so that conservation efforts that seek the implementation of new marine protection areas can have reliable data so as not to underestimate
27 the biodiversity of species (Sala et al., 2021).

In the same way, by overlapping the bioregions with the fishing exploitation zones, we determined that the North Pacific (7) North West Pacific (29),
30 Mid-tropical N Pacific Ocean (9) and Indo-Pacific seas and Indian Ocean (13)

bioregions , as well the Gulf of California (21) and Caribbean and Gulf of Mexico (11) are the regions with the highest representation of the data and 3 where fishing activity is concentrated. According to ([Kroodsma et al., 2018](#)), the area corresponding to the central Atlantic and Northeast Pacific present little intense fishing effort, while the regions associated with the Northeast 6 Atlantic, the Northeast Atlantic (Europe) regions, and the Northwest Pacific are known to have a huge fishing development and that is where fishing efforts are concentrated worldwide. The southeastern Atlantic Ocean (FAO 9 area 47 and 88), part of the Pacific Ocean (FAO area 88) and Antarctica (FAO area 48 and 88) are the regions with the highest percentage of cells without records ($NR = >93\%$). When compared with the findings of ([Kroodsma 12 et al., 2018](#)), these areas agree with the “holes” without fishing effort data, which is explained by the geographical remoteness and the lack of technological development necessary for the fisheries to extend to new domains ([Visalli 15 et al., 2020](#)). This limits both the exploitation of marine resources and the collection of data.

The hypotheses that we evaluated in this work were necessary to understand 18 what the data collection trends have been and to be able to take future actions to correct the biases described. Our first hypothesis about the size of the body of the fish was rejected. Small fish species (0-80 cm) are the ones 21 that accumulate the largest number of records, among which *S. scombrus* (Scombridae), *L. rhomboides* (Sparidae), *M. villosus* (Osmeridae), are <50 cm species that stand out for presenting the largest number of records, and, 24 in addition, they are distributed in the best sampled regions (Mediterranean Sea, Gulf of Mexico and the Caribbean, and the Atlantic Ocean). The size of the fish is inversely proportional to the abundance and, therefore, to the frequency 27 of human use, both scientific and commercial ([Pauly and Palomares, 2005](#)). This difference in the sampling effort generates an evident overrepresentation of the smaller species and therefore deepens the taxonomic bias. 30 The hypothesis about the depth of the habitat is accepted, at less depth

there is a greater representation of species of marine fish. The pelagic zone has a high concentration of data and effectively corresponds to shallow regions and therefore easily accessible, which generates all the conditions for data collection (Melo-Merino et al., 2020). It has been pointed out that the concentration of species decreases as the depth of the ocean increases, however, it is precisely these areas that have been least sampled and where there is the greatest probability of discovering new species (Costello et al., 2017). This demonstrates the need to concentrate efforts on the deeper regions of the water column (mesopelagic, bathyal, and abyssal) for a more equitable representation of marine ecosystems. Finally, the hypothesis of the use of the species is also accepted. The species of marine fish that have a more beneficial or lucrative use for humans are better represented in the analyzed databases. We believe that this is related to the fact that the fishing industry is one of the main sources of information for platforms such as OBIS (Zhang and Grassle, 2002).

Today, marine ecosystems and their biodiversity face the great challenges of climate change and the impact of human activity, especially those species considered key food resources for survival (Hollowed et al., 2013; Ramírez et al., 2017; O'Hara et al., 2021). It is necessary to focus and strengthen the study of those areas with very few or no records, since the descriptions of the geographic ranges of the species and their temporal dynamics are fundamental measures for the evaluation of the real state of biodiversity (Lenoir et al., 2020; Oliver et al., 2021). Having more reliable data will allow effective conservation actions to be implemented.

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Appendix A. The database

Table A.4 below shows the data loss for each criterion that we have used
3 to clean our database. We downloaded 71,670,596 records from GBIF and OBIS. Only 820,004 records were useful for our analyses.

Database state	Number of records
Origial records from GBIF and OBIS	71,670,596
Data curation (following Zizka et al. (2020))	5,380,439
Taxonomically filtered data	5,007,322
Deletion of data outside the native range	820,004

Table A.4: Criteria for filtering occurrence data from GBIF and OBIS using bioregions.

Files of the 10,371 marine fish species and their attributes (body size,
6 habitat depth, and cultural value) from FishBase may be found in the GitHub project page of this manuscript: http://github.com/vapizarro/stp_fishes

Appendix B. Grids resolutions

9 For spatial representation analysis we evaluated two additional spatial resolutions ($5^\circ \times 5^\circ = 3,021$ cells, and $10^\circ \times 10^\circ = 958$ cells). Table B.5 contains the results of this analysis for these grids. We have also mapped these
12 results (see Figure B.7), to understand how the effect of spatial resolution on the evaluation of biodiversity macropatterns. Finally, we also plot the frequency of cells for each SRI category for the three grid sizes (R1= $1^\circ \times 1^\circ$;
15 R5= $5^\circ \times 5^\circ$; R10= $10^\circ \times 10^\circ$) to understand how the data is distributed in our analyses (see Figure B.8)

Appendix C. Bioregions slopes

18 We evaluated the slopes of the last 10% of the accumulation curves of each bioregion in our temporal representation analysis. Table C.6 shows the result for each bioregion.

ID	R1 ($1^\circ \times 1^\circ$)					R5 ($5^\circ \times 5^\circ$)					R10 ($10^\circ \times 10^\circ$)				
	NR	IR	F	S	A	NR	IR	F	S	A	NR	IR	F	S	A
1	18.49	15.13	5.04	36.97	24.37	0.00	16.67	0.00	33.33	50.00	16.67	16.67	0.00	33.33	33.33
2	68.75	19.79	1.04	10.42	0.00	10.00	40.00	10.00	30.00	10.00	40.00	0.00	0.00	40.00	20.00
3	15.74	6.54	3.39	36.80	37.53	3.57	3.37	0.00	17.86	75.00	0.00	0.00	0.00	10.00	90.00
4	46.35	22.34	7.93	22.13	1.25	28.13	9.38	6.25	40.63	15.63	30.77	15.38	0.00	23.08	30.77
5	42.39	14.75	4.92	27.87	10.07	14.29	3.57	0.00	32.14	50.00	16.67	8.33	0.00	8.33	66.67
6	94.96	2.21	0.13	1.87	0.83	82.13	5.64	0.31	6.58	5.33	62.65	13.25	1.20	10.84	12.05
7	63.24	11.24	0.87	14.46	10.19	17.09	9.40	3.42	29.06	41.03	7.69	7.69	2.56	35.90	46.15
8	79.52	11.27	0.89	7.17	1.15	43.93	11.56	4.05	32.37	8.09	32.69	11.54	3.85	40.38	11.54
9	88.74	8.71	0.00	1.57	0.99	28.74	22.99	2.87	38.51	6.90	7.69	9.62	21.15	38.08	13.46
10	96.31	2.41	0.04	0.88	0.36	70.87	15.75	0.79	7.09	5.51	51.28	15.38	2.56	20.51	10.26
11	23.82	8.42	5.65	32.85	29.26	8.62	0.00	0.00	18.97	72.41	0.00	10.53	0.00	5.26	84.21
12	35.59	21.61	2.45	35.59	4.76	14.29	4.76	2.38	47.62	30.95	5.88	11.76	0.00	17.65	64.71
13	67.52	15.80	1.01	12.00	3.67	13.76	12.84	7.34	44.95	21.10	9.46	6.76	2.70	44.59	36.49
14	45.83	10.83	2.50	30.83	10.00	46.15	0.00	0.00	7.69	46.15	25.00	0.00	0.00	0.00	75.00
15	74.52	13.06	0.00	7.07	5.35	20.00	6.67	6.67	40.00	26.67	37.50	12.50	0.00	37.50	12.50
16	36.68	10.95	3.84	34.65	13.88	5.77	7.69	3.85	28.85	53.85	10.53	0.00	0.00	21.05	68.42
17	91.36	4.90	0.00	1.57	2.17	47.93	19.01	0.00	20.66	12.40	25.00	8.33	0.00	36.11	30.56
18	48.29	16.27	3.78	22.06	9.61	6.50	7.32	7.32	43.09	35.77	10.26	5.13	0.00	28.21	56.41
19	90.40	6.93	0.06	2.27	0.35	53.45	18.39	3.45	17.82	6.90	31.48	12.96	1.85	35.19	18.52
20	63.61	17.35	1.43	13.56	4.04	8.20	8.20	9.84	44.26	29.51	15.00	5.00	0.00	45.00	35.00
21	74.78	9.63	2.84	11.48	1.27	34.68	13.51	3.15	28.38	20.27	21.21	9.09	0.00	27.27	42.42
22	76.12	18.00	0.00	5.10	0.78	33.33	6.17	16.05	43.21	1.23	9.09	4.55	9.09	59.09	18.18
23	34.65	32.02	2.89	24.41	6.04	25.93	0.00	14.81	51.85	7.41	0.00	18.18	0.00	36.36	45.45
24	63.07	17.89	1.38	13.53	4.13	25.93	7.41	3.70	51.85	11.11	20.00	10.00	0.00	50.00	20.00
25	88.02	4.96	0.83	5.37	0.83	52.63	10.53	0.00	21.05	15.79	42.86	0.00	0.00	28.57	28.57
26	60.93	7.04	2.41	22.41	7.22	27.27	12.12	0.00	30.30	30.30	16.67	0.00	0.00	33.33	50.00
27	66.84	10.35	0.70	8.07	14.04	27.78	16.67	0.00	22.22	33.33	7.69	7.69	0.00	23.08	61.54
28	59.84	9.17	2.13	17.67	11.19	30.19	7.55	1.89	30.19	30.19	30.00	10.00	0.00	25.00	35.00
29	41.96	19.87	2.84	26.81	8.52	15.00	10.00	5.00	40.00	30.00	12.50	12.50	0.00	37.50	37.50
30	93.74	3.49	0.20	1.97	0.59	69.45	11.02	1.00	10.52	8.01	42.29	16.57	2.86	22.29	16.00

Table B.5: Surface area as a percentage of each bioregion (ID) for every SRI category for each of the three grid sizes (R1= $1^\circ \times 1^\circ$; R5= $5^\circ \times 5^\circ$; R10= $10^\circ \times 10^\circ$). Values show the surface area as a percentage of each bioregion for every SRI category (see §2.2.1). ID is the identification number given to each bioregion (Table 1). A are cells with an adequate representativeness of species richness (i.e. SRI > 0.85). S are cells considered as having a sufficient representativeness (i.e. SRI $\in (0.60, 0.85)$). F cells are cells with few records and are thus not considered to be representative of actual species richness (i.e. SRI $\in (0, 0.6)$). NR as cells with no records (SRI= NA), and IR as cell with insufficient records to apply SRI.

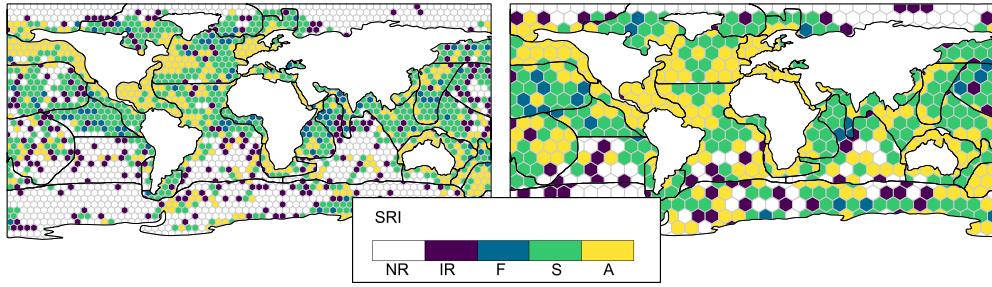


Fig. B.7: Spatial representativeness index (SRI) mapping of cells of size: $A=5^\circ \times 5^\circ$; $B=10^\circ \times 10^\circ$. The categorization of the cells corresponds to the level reached by the SRI, where $SRI > 0.85$: Amount of data *Adequate* for the representation of species richness (“A”); $SRI=0.60-0.85$: Amount of data can be considered *Sufficient* (“S”); $SRI=0-0.60$: Amount of records *Few* (“F”); and $SRI = NA$: cells with no records (“NR”).

Appendix D. Supplementary Material: GAP Analysis

We plotted the percentage of surface with marine protected areas of each
³ bioregion (Fig D.9), and the percentage of cells of each FAO Area for each category of SRI value (Fig D.10).

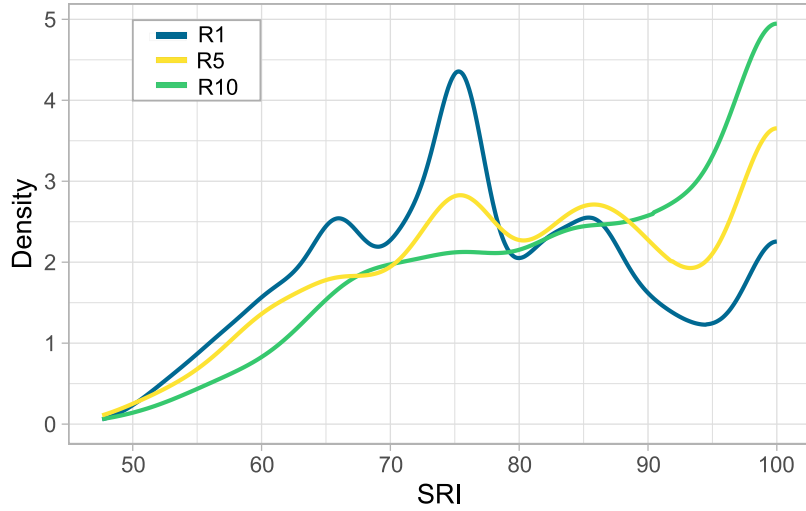


Fig. B.8: Density probability distribution of SRI in three grids of different sizes: R1 = $1^\circ \times 1^\circ$ (blue line); R5 = $5^\circ \times 5^\circ$ (red line); and R10 = $10^\circ \times 10^\circ$ (yellow line).

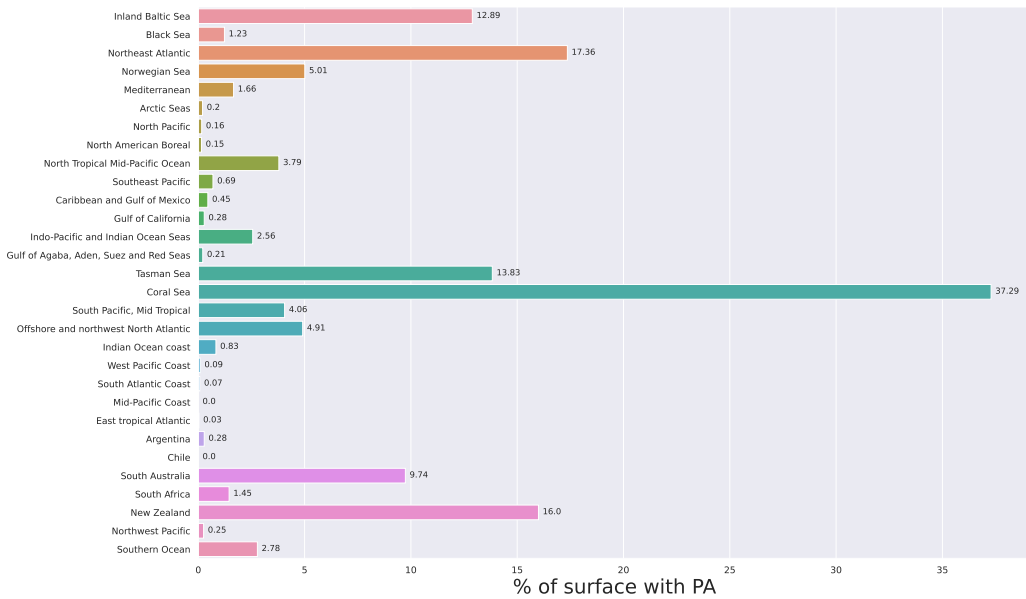


Fig. D.9: Percentage of surface with marine protected areas by bioregions.

Bioregion	Slope
1	0.35
2	1.16
3	1.79
4	0.91
5	1.76
6	0.65
7	4.44
8	1.37
9	6.18
10	4.87
11	10.37
12	7.57
13	32.86
14	4.90
15	6.78
16	21.62
17	10.10
18	6.59
19	12.44
20	23.21
21	11.70
22	1.85
23	4.42
24	3.49
25	2.12
26	7.74
27	12.29
28	4.82
29	14.08
30	2.74

Table C.6: Final slope (10%) of the accumulation curves for each bioregion

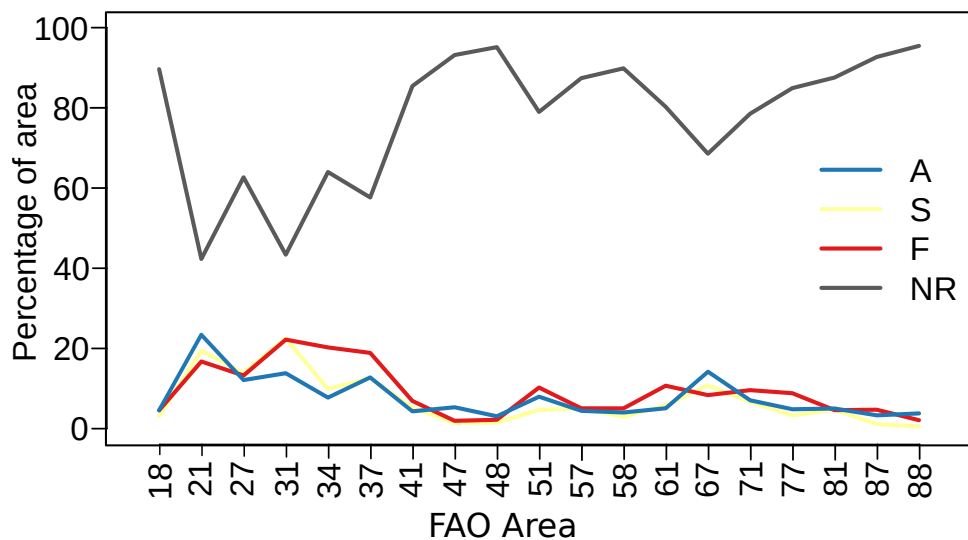


Fig. D.10: Percentage of cells of each FAO Area for each category of SRI value. Amount of data *Adequate* for the representation of species richness (“A”); SRI=0.60-0.85: Amount of data can be considered *Sufficient* (“S”); SRI=0-0.60: Amount of records *Few* (“F”); and SRI = NA: cells with no records (“NR”).