1 2	Influence of Oxygen Minimum Zone on Macrobenthic Community Structure in the Northern Benguela Upwelling System: A Macro-Nematode Perspective
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Abstract

for predicting the impacts of climate change and human activities on oceanic biodiversity. In this 13 context, we conducted a study on the macrobenthic communities in the Benguela Upwelling 14 System (BUS). Macrobenthic samples were collected offshore Namibia on board R/V Mirabilis 15 16 during the 3rd RGNO training and the National Marine Information and Research Centre's (NatMIRC's) plankton survey from 13th May to 17th May 2016. Two transects, Terrace Bay, (20° 17 S) and Walvis Bay (23° S), hosted three stations each, while the third transect, Luderitz (26° S), 18 19 hosted only one station. From the results, three oxygen zones were identified, namely Microxic (<0.1 ml l⁻¹), Dysoxic (0.1-1.0 ml l⁻¹), and Oxic (>1.0 ml l⁻¹). A total of 20 Macrobenthic taxa were 20 21 identified; Nematoda, Polychaeta, Cumacea, and Oligochaeta were the most dominant taxa and 22 were present in all oxygen zones. Eighteen genera of macro-nematoda were identified; Desmolaimus and Paracomesoma dominated in all oxygen zones, Metoncholaimus recorded 23 higher abundance in the dysoxic stations, and were absent in the oxic stations, and the opposite 24 was observed for Halanonchus and Dorylaimopsis. H' Diversity values for both the general 25 26 macrofauna and macro-nematoda were higher in the oxic stations and lower in both the dysoxic and microxic stations, while an opposite trend was observed for dominance values. Density values 27 were lower in microxic stations and higher in dysoxic stations, while at the oxic stations, the 28 density values fell in between the two hypoxic zones. In conclusion, this study provides an 29 overview on the distribution, diversity, and response to varying oxygen conditions of 30 macrobenthic communities and their importance in marine ecosystems. 31

Understanding the dynamics of marine ecosystems, particularly the benthic communities, is crucial

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51 Keywords : Macrofauna, Macro-Nematoda, Oxygen minimum zone, Benguela Upwelling Sy

52 (BUS), Dissolved oxygen, hypoxia

1.0 Introduction

In recent decades, the concentration of dissolved oxygen (DO) in the ocean, specifically in the tropics, has been decreasing. This not only increases the size of areas under hypoxia but also their prevalence (Breitburg et al., 2018). Hypoxia is ranked among the major threats to the actualization of the blue economy and the achievement of the 14th pillar of the Sustainable Development Goals (SDGs), specifically target 14.7 which aims at increasing the economic benefits of marine resource utilization through its sustainable use by developing countries (Rossi, 2023; Liu, 2023). Although hypoxia can result from natural phenomena like upwelling and thermal stratification, the current expansion of hypoxic areas is mainly a result of accelerated nutrient inputs in coastal areas which increases algal production subsequently resulting in higher organic matter production which in turn results in increased aerobic microbial decomposition lowering the levels of DO in the water (Gobler & Baumann, 2016). It has been projected that such changes may affect different organisms differently depending on their tolerance and reactions to lower dissolved oxygen in their habitats (Rudolph et al., 2020). Studies on ecosystems with hypoxia as a natural phenomenon can assist in predicting and understanding how human-induced hypoxia might affect and shape marine ecosystems in the face of the expansion of marine areas under hypoxia.

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The Benguela upwelling system (BUS) is located off the southwest coast of Africa. It extends from

Cape Frio in Angola to the southern tip of the continent in Cape Agulhas, South Africa, and is one

of the most productive regions of the world's ocean (Magalhães, 2018). The high productivity

provides a huge source of carbon resulting from photosynthesis, which gradually sinks through the

water column resulting in a rapid oxygen loss due to biochemical oxygen consumption. This consequently causes low dissolved oxygen concentrations and thus creates a permanent extensive shallow oxygen minimum zone (Bohata & Koppelmann, 2013; Emeis et al., 2018). When the oxygen minimum zone (OMZ) comes into contact with the seafloor, it creates a strong oxygen gradient at the benthic zone at depths ranging between 50 meters and 300 meters, resulting in a hypoxic (<0.5 ml 1⁻¹) inner shelf (Gibson & Atkinson, 2003). The oxygen gradient created at the OMZ's benthic zone is believed to primarily regulate the benthic community distribution and diversity patterns (Zettler et al., 2013; Teuber et al., 2013). In these zones, microbial communities play a key role in the cycling of elements like carbon, nitrogen, and sulfur. The expansion of OMZs may increase nitrate use by nematode associated microbes such as SAR11, a significant player in the marine nitrogen cycle. These nematodes and the communities they are part of influence biogeochemical processes and ecosystem dynamics within OMZs. For this reason, understanding these communities is crucial, especially with the ongoing expansion of OMZs due to climate change.

The general trend observed in most OMZs in global oceans namely, Walvis Bay, Namibia (the location of this study), California, USA and Oman margin (off the Arabian Peninsula) indicates that the densities of macrofauna (benthic organisms large enough to be seen with the naked eye and are typically retained in a 0.5 mm sieve but pass through a 2.00 mm sieve (Bachelet, 1990), generally display a negative response to reducing oxygen levels within the OMZ with a 30% to 70% reduction in densities in regions with less than 0.15 ml l⁻¹ (Gibson & Atkinson, 2003). Similarly, diversity reduces as oxygen levels reduce within the OMZ because of the loss of intolerant species and increased dominance of the tolerant species (Currie et al., 2018; Levin et al., 2009). Nematodes and some families from the annelid worms have been observed to be able to

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Deleted: The term' benthos' refers to organisms living on and in the sediments of the seafloor and are distinguished based on their sizes as either megafauna, macrofauna, meiofauna, or microfauna with some taxa appearing in more than one size category. Macrobenthos are part of the benthos consisting of organisms retained in a 0.5mm sieve but pass through a 2.00mm sieve (Bachelet, 1990).

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112 tolerate low oxygen, with Nematoda (in the meiofauna group) having 95-99% abundance. Some polychaetes families like Spinoid, Dorvilleid, and Lumbrinerid can also tolerate low oxygen in the 113 OMZ by having a high gill surface area for increasing oxygen uptake. On the other hand, 114 115 harpacticoid copepods are the most sensitive taxon to hypoxia (Levin et al., 2009; Zeppilli et al., 2015). 116 117 It has been observed that the macrofauna diversity is lowest off Walvis Bay, a city located on the western coast of Namibia, attributed to the perennial intense OMZ over the shelf (Currie et al., 118 119 2018). In contrast, the diversity increases significantly northwards off the Kunene River (Zettler 120 et al., 2013), which flows from the highlands of Angola, along the border with Namibia, and into 121 the Atlantic Ocean (See Figure 1). This increase in diversity is proposed to be a result of moving 122 away from the intense OMZ cells off Walvis Bay and also the reflection of the biogeography where 123 diversity increases with reducing latitude (Zettler et al., 2009). Outside the OMZ, bathymetry, and 124 latitude are said to be the factors affecting the infaunal communities at the Namibian shelf (Steffani 125 et al., 2015). This study aims to identify the relationship between the levels of dissolved oxygen and the 126 macrobenthic community distribution across the Northern Benguela Upwelling system as a guide 127 128 on how the expansion of man-made hypoxia might influence the benthic fauna distribution on the seafloor. 129 130 2.0 Material and Methods 131 2.1 Study Area The study area was located across the Northern Benguela Upwelling System (between 26° S and 132 20° S) along the Namibian continental shelf, which hosts a deep continental shelf (around 300 m)

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137 (Figure 1). The intense upwelling in the study site has rendered the area highly productive, Commented [TM12]: Reference to Figure 1 added at end of sentence. resulting in a characteristic shallow OMZ (50-200 m) with stronger productive cells around 138 Luderitz and Walvis Bay (Bohata & Koppelmann, 2013). The inner shelf is described to be 139 140 extremely oxygen-depleted, caused by in situ organic matter decomposition and warm poleward Angola current, which peaks in June-July while the continental slope below the OMZ is better 141 oxygenated (Levin et al., 2009; Emeis et al., 2004). 142 The benthic zone in the OMZ in Northern BUS is characterized by extensive areas of diatomaceous 143 144 mud, which are associated with high primary production at the ocean surface and low Commented [TM13]: Specified to ocean surface Deleted: surface concentration of dissolved oxygen (Levin et al., 2009; Steffani et al., 2015). These low oxygen 145 levels, as per the findings of Levin et al. (2009) were found to be less than 0.5 ml L-1 in 55% of 146 147 the total shelf. Furthermore, extreme anoxia, defined as oxygen concentrations less than 1 µM, was observed over an expansive area of almost 900 km². 148 Commented [TM14]: This information has been added to clarify how low the oxygen levels were 149 2.1 Sample Collection Samples were collected from three transects (off Luderitz (OL) 26° S, off Walvis Bay (OWB) 23° 150 Deleted: Cape Frio 151 S, and off Terrace Bay (OTB) 20° S) onboard R/V Mirabilis during the RGNO training National Deleted: CF Marine Information and Research Centre's (NatMIRC's) plankton survey from 13th May to 17th 152 153 May 2016. The sampling stations were located at 02 nm, 20 nm, 40 nm, or 70 nm at each transect, 154 with the 26° S transect hosting only one station at 90 nm (Figure 1). However, benthic samples 155 from these stations were dependent on the prevailing weather conditions and the ability to get good 156 core samples (Table 1). Commented [TM15]: Figure 1 changed to Table 1 Deleted: Figure 157 Sampling was done using a multi-corer from which individual cores were taken and sub-sampled 158 for macrobenthos using a 6.4 cm diameter corer, and sediment samples for granulometry and

organic matter analysis were taken simultaneously. Replicate samples were taken from a subsequent deployment of the multi-corer, where possible, to avoid pseudoreplication. The macrofauna cored samples were put in sampling bottles and immediately fixed with 5% buffered formalin, while samples for organic matter analysis were frozen to arrest microbial decomposition. Depth and temperature measurements were collected from a probe attached to the multi-corer, while dissolved oxygen concentrations in this study used the Winkler method from the overlying water (Montgomery et al., 1964).

2.3 Laboratory analysis

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In the laboratory, macrobenthic samples were sieved between 2.00 mm and 0.45 mm_sieves.

Studies typically employ a sieve size of 0.5mm, our study however utilized a 0.45mm sieve due to its availability during the course of our research. This size, although slightly smaller, falls within the range of commonly used sieve sizes (300 to 500 microns). The use of a 0.45mm sieve is also not unprecedented in macrofaunal studies; studies by Li et al. (2018) and Zhang et al. (2022) have successfully employed this sieve size. Therefore, while acknowledging the slight deviation, results remain valid and comparable to those obtained using the more common sieve sizes.

The sieve fraction retained in the 0.45 mm mesh sieve was then preserved in 5% buffered formalin solution, with 3-5 drops of Rose Bengal solution added to aid in sorting. After rinsing the samples with water, they were sorted under a dissecting microscope using the key provided by Thiel & Higgins (1988) to the class or order levels, and then counted. Subsequently, the nematodes were pin-picked, fixed on permanent slides, and identified to the genus level using the key from Platt & Warwick (1988). Feeding types were determined for these genera following the methodology described by Wieser (1953), whereby nematodes are classified based on their buccal morphology,

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including shape, size, and presence or absence of buccal armature, resulting in four groups. Those lacking buccal armature are categorized as deposit feeders (group 1), further subdivided into selective deposit feeders (1A) and non-selective deposit feeders (1B) based on the size of their buccal cavity, with the former possessing a very small buccal cavity, and the latter having a larger one (Moens and Vincx, 1997; Moens et al., 2013). Nematodes with buccal armature constitute group 2, and can be Selective epigrowth feeders and herbivores (2A), using small teeth to graze diatoms, microalgae, and bacteria, or carnivores and omnivores (2B), characterized by the possession of teeth, mandibles, or onchia for piercing and consuming prey (Moens et al., 2013) For sediment granulometry, samples were wet-sieved using a 1.00 mm sieve, 0.5 mm, 0.25 mm, 105 mm, 0.063 mm, and those below 0.063 mm (<0.063 mm). The sediments collected at the different sieves were then put on pre-weighed Petri dishes and dried in an oven at 60° C until a constant weight was reached, and these dried weights were used for sediment granulometry (Wentworth, 1922). The samples for total organic matter (TOM) were dried in an oven at 60° C until a constant weight was achieved, then a known weight of each sample was then placed in labeled aluminum foil, and the loss of weight on ignition (LOI) technique was used to determine TOM from the study site

2.4 Data analysis

(Thiel & Higgins, 1988).

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Out of the seven sampled stations, only three had replicates (Table 1), making direct statistical comparisons between stations impossible. To overcome this limitation, the sampling stations were grouped based on the measured environmental variable that provided the most significant relationship identified through step-wise regression analysis with the various biotic indices

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233 were then compared using the indices and analyses outlined below; Deleted: was used to group the stations for proper community analysis. Grouping Stations; Following the approach of (Levin, 2003), stations with DO levels less than 0.1 234 Commented [TM18]: Reference now reads (Levin, 2003) ml 1⁻¹ were grouped as microxic. those with DO levels between 0.1 and 1.0 ml 1⁻¹ were grouped 235 Commented [TM19]: Microxic as per suggestion not to 236 as dysoxic and those with DO levels of 1.0 ml l⁻¹ and above grouped as oxic. This approach was Commented [TM20]: dysoxic 237 taken to investigate the role of DO levels in these communities. It was further justified by the Commented [TM21]: oxic Deleted: 238 application of SPSS, which indicated that DO was a key determinant in macrobenthic community Deleted: so as structures ($R^2=0.521$, P=.012) (Table S1a; S1b(1); S1b(2)). 239 Formatted: Font: Italic Abundance Analysis; The data recorded were analyzed for abundance (density and relative 240 Deleted: The data recorded 241 abundance) in Excel. Additionally, Paleontological Statistics Software package (PAST v2.17c) Deleted: (Hammer et al., 2001) was used to calculate the diversity and the diversity t-test between the 242 243 stations. Community Similarity and Diversity; Bray Curtis similarity, Analysis of Similarity (ANOSIM), 244 Commented [TM23]: As youll probably notice I adopted the alternative approach, which included stating and justifying that weiser's method could be used for analysis. a 245 and Similarity percentages (SIMPER) were conducted using Plymouth Routines in Multivariate fact that was justified by spss showing do as a vital value in macrobenthic communities. This part was a bit tricky for Ecological Research (PRIMER v5.2.9) (Clarke & Gorley, 2005). 246 me, but I hope it fit the criterion by both reviewers. Deleted: Predictive Modelling; Stepwise regression analysis (using SPSS) was used to determine the 247 3.0 Results environmental variables that were predictors of the distribution patterns of the macrobenthic diversity patterns in the Benguella Upwelling USystem. Various diversity indices (considering replicates as a single station) indices were 248 3.1 Abiotic variables calculated using PAST for each station; the various replicates were considered as a single station, and these indices were then correlated individually with the environmental variables recorded in the study.¶ 249 Total Organic Matter (%TOM) demonstrated an inverse relationship with depth, with higher Commented [TM24]: Inverse relationship with inserted 250 organic matter values recorded in the shallower stations. For example, the shallowest stations Deleted: showed an opposite trend Commented [TM25]: Sentence structure altered 20020, 20002, and 23002 had the highest TOM of 38.6% ±2.16, 27.5% ±1.55 and 23.1%, 251 Deleted: where

documented within the macrofauna community. The communities within these key 'habitat types'

respectively whilst the lowest TOM values were found in most offshore stations, stations 23070

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and 26090, with 9.89±1.55 and 4.4, respectively (Table 1). The lowest oxygen values were recorded on the OWB 23° S transect, with the two most onshore stations (23020 and 23002) recording the lowest levels of oxygen (0.02 and 0.06 ml l⁻¹, respectively). The inshore station from transect OCF 20° S (20002) had a higher DO (1.8 ml l-1) than stations 20020 and 20040 from the same transect, where the DO levels recorded were 0.67 and 0.53 ml l⁻¹, respectively, while the offshore stations 23070 in transect OWB 23° S and 26090 in transect OL 26° S recorded the highest dissolved oxygen (2.30 and 4.34 ml l⁻¹, respectively) (Table 1).

Fine sand was the most common sediment size ranging between 30-38.8% in all stations. Most stations from the sites also recorded higher proportions of medium sand and silt, except for station 23020 in transect OWB 23⁰ S, which recorded a lower proportion of silt and an increased abundance of coarse sand compared to the other stations (Table 2).

3.2 Macrobenthic assemblages

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Macrofaunal densities differed significantly across the various oxygen zones; <u>In the microxic stations</u>, very low densities were observed (4,661±4,834 ind. m₂⁻²), and the lowest number of taxa was recorded, with only six taxa present. These included Nematoda, Polychaeta, Oligochaeta, and Cumacea, the only peracarid crustacean found across all oxygen zones. Despite the low number of taxa (S=6), these stations recorded slightly higher diversity indices than the dysoxic stations, while the dominance index was intermediate (0.5±0.18) (Figure 5).

station within this Oxygen range, station (20040), recorded an extremely high density of 274,991 ind. m-2, primarily due to a high abundance of Nematoda and Oligochaeta. Although the taxa in the dysoxic stations was similar to that of the microxic stations, the composition and dominance

Dysoxic stations contrastingly recorded the highest densities (74,108±134,126 ind. m-2). One

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Stepwise regression analysis revealed that most diversity indices (H' Diversity, Evenness, Dominance, Berger-Parker, Fisher Alpha, Equitability J, and Brillouin) had no predictive variables except Margalef index, Richness, and Menhinick whose predictors were identified as TOM, DO, and very coarse sand respectively. Pearson correlation analysis indicated a significantly high correlation between DO and TOM, as TOM reflects surface production, which is the driver of the low DO at the BUS (Table S1). ¶ Due to the lack of replicates in some stations and the high predictive role in DO compared to the other two factors, the sampling stations were thus grouped based on their recorded levels of DO as described by (Levin Lisa, 2003). i.e., stations with DO less than 0.1 ml 1-1 were grouped as 'Microxic.' those with DO between 0.1 and 1.0 ml l-1 grouped as 'Dysoxic' while those with DO of 1.0 ml l-1 and above grouped as 'Oxic' in a bid to analyze how DO affects macrobenthic communities structure.

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20	differed gianificantly. The moore nemated expectly most dominant toys in this zone, with numbers		
30	differed significantly. The macro-nematoda was the most dominant taxa in this zone, with numbers	/	Deleted: A total of 19 macrobenthic taxa were identified and were dominated by Nematoda, Polychaeta, Oligoc
31	reaching as high as 233,354 ind. m-2 in one of the stations. Interestingly, Bivalvia and Ostracoda,		and Cumacea, the only peracarid crustacean abundant
32	absent in the microxic stations, were present here, albeit in low abundance (Figure 3). Dominance		the oxygen zones. Microxic stations recorded the lowe count (S=6), which constituted Cumacea (37.5%), Polychaeta (26.9%), Nematoda, Oligochaeta, and Asci the only taxa present in the Microxic stations (Fig 3).¶
33	was highest in these stations (0.6 ± 0.28) (Figure 5).	/	Deleted: Although the taxa count in the dysoxic static
34	Oxic stations demonstrated the highest number of taxa (18) and the second-highest average density (14,345±6,726 ind. m-2) (Figure. 2). These stations recorded taxa like Echinodermata		was like that of the microxic stations, the composition dominance were different. The macro-nematoda was the most dominant taxa in this zone, outnumbering all the taxa. Its numbers were very high in one of the stations (233,354 ind. m ²). Oligochaeta was the second most abundant taxon in this oxygen zone, with its high abundant taxon in this oxygen zone, with its high abundant taxon in the second most abundant taxon in this oxygen zone.
36	(Ophiuroidea), Holothuroidea, Aeolosomatidea, Isopoda, Aplacophora, and Amphipoda, which all		coming from the same station with the highest nemato counts. Polychaeta and Cumacea were the dysoxic stat third and fourth most abundant taxa. Bivalvia and Ostr which were absent in the microxic stations, were prese the dysoxic stations, albeit in low abundance (Figure 3
37	had average abundances of more than 1%. Nemertina, Turbellaria, and Cnidaria were also recorded		Deleted: The oxic stations recorded the highest numb
38	in these stations; however, their abundances were below 1%, and they were grouped as 'others'		taxa counts, with 18 out of the 19 taxa recorded overal like Echinodermata (Ophiuroidea), Holothuroidea, Acolosomatidea, Isopoda, Aplacophora, and Amphipo
39	(Figure 3). These stations recorded the highest H' diversity (1.46±0.4) and the lowest dominance		were recorded in this zone with average abundances of than 1%. Nemertina, Turbellaria, and Cnidaria were al recorded in these stations; however, their abundances
40	(0.31±0.18) (Figure 5),		below 1%, and they were grouped as others (Figure 3)
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42	using Bray-Curtis analysis of dissimilarity. The highest dissimilarity was observed between the	////	Deleted: M
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43	dysoxic and microxic zones which were 77.99% dissimilar despite both stations being	' / // _/	Commented [TM42]: This entire segment has been
344	characterized by low dissolved oxygen levels. The oxic stations were also highly dissimilar to the		altered to make way for reviewer 2's suggestion that a station should have its own paragraph. I modified it to so even the original flow has some formats according reviewers suggestion.
45	microxic and dysoxic zones, with values of 68.58% and 65.91%, respectively (Figure 4).		Deleted: The Dysoxic and Microxic stations recorded macrobenthic H' diversity (<1), with the Microxic station
46	3.3 Macro-nematodes density and diversity	///	Deleted: ¶
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47	On average, macro-nematodes were the most dominant taxon in this study as a result of their		Commented [TM43]: Just nematodes after first me - reviewer 2
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48	dominance in the dysoxic station. Nematodes were abundant in all oxygen zones recording relative		Deleted: M
49	abundances of 8%, 74%, and 24% in microxic, dysoxic, and oxic zones respectively (See Figure		Deleted: D
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410	In our study, eighteen different genera of macro-nematodes were identified. Notably,
411	Desmolaimus and Paracomesoma were detected across all stations, and were the only genera
412	present in microxic stations. These two, along with Metoncholaimus, exhibited their highest
413	abundance in dysoxic stations. Additionally, in these dysoxic environments,
414	Paralongicyatholaimus and Neochromadora recorded high abundances, each constituting (>4%)
415	of the total population Figure 6). Thalassolaimus, Paramesacanthion, Enoploides, Halanonchus,
416	Rhabdodemania, and Dorylaimopsis recorded high abundances in oxic stations but were absent in
417	dysoxic stations except for Thalassolaimus. Metoncholaimus and Paralongicyatholaimus, were
418	present in dysoxic stations but absent in oxic stations, while Paramesacanthion, Enoploides, and
419	Rhabdodemania were present in the oxic station and absent in the dysoxic stations. For the
420	purposes of graphing the relative abundance, Thoracostomopsis, Anticoma, Cephalanticoma,
	Trileptium, Mesacanthoides, Terschellingia, and Marylinnia were grouped as 'others' as they
421	Truepium, mesacuunoues, Terscheungur, and maryumuu were grouped as Johiers_ as they
421 422	recorded <u>low</u> abundances (<4%) and were absent in <u>dysoxic stations</u> except for <i>Marylinnia</i> and
422	recorded low abundances (<4%) and were absent in dysoxic stations except for Marylinnia and
422 423 424	recorded <u>low</u> abundances (<4%) and were absent in <u>dysoxic stations</u> except for <i>Marylinnia</i> and <i>Terschellingia</i> , whereby, the former was absent in the <u>oxic station</u> while the latter was present in both oxygen zones (<u>See Figure 6</u>).
422 423 424 425	recorded <u>low</u> abundances (<4%) and were absent in <u>dysoxic stations</u> except for <i>Marylinnia</i> and <i>Terschellingia</i> , whereby, the former was absent in the <u>oxic station</u> while the latter was present in both oxygen zones (<u>See Figure 6</u>). The feeding guild composition exhibited variation across distinct oxygen zones within the study
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422 423 424 425 426 427 428 429	recorded low abundances (<4%) and were absent in dysoxic stations except for <i>Marylinnia</i> and <i>Terschellingia</i> , whereby, the former was absent in the oxic station while the latter was present in both oxygen zones (See Figure 6). The feeding guild composition exhibited variation across distinct oxygen zones within the study site. Selective epigrowth feeders and herbivores, categorized as Type 2A, emerged as the predominant nematodes within dysoxic zones, comprising 40% of the population. This was closely followed by non-selective deposit feeders (Type 1B), contributing to 35% of nematodes within the same zone. Carnivores and omnivores, classified as Type 2B, constituted 28% of the nematode

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nematodes, respectively. Carnivores and omnivores (Type 2B) and selective deposit feeders (Type 1A) accounted for 18% and 15%, respectively (Figure 7). Within the feeding modes, selective deposit feeders had the largest differences in composition between the two oxygen zones, with the highest concentration observed in the oxic zones. While 2A and 2B had slightly higher values in the dysoxic zone compared to the oxic zones.

The nematode diversity exhibited a pattern akin to the overall macrofaunal diversity, wherein one of the microxic stations (23002) was devoid of nematodes, while its counterpart station (23020) harbored only two nematodes. Consequently, station 23002 was excluded from diversity analysis. Across the remaining oxygen ranges (dysoxic and oxic), a parallel trend was observed as in the broader macrofaunal analysis within the study area. Specifically, oxic stations displayed higher H' Diversity (1.38±0.5) compared to dysoxic stations (0.81±0.84). Conversely, dysoxic stations exhibited greater dominance (0.59±0.39) relative to oxic stations (0.32±0.18) (Fig. 8).

4.0 Discussion

Upwelling systems are known for their high surface productivity and Oxygen Minimum Zones (OMZ), which impinge on the benthic zone creating strong oxygen gradients on the seafloor and acting as the dominant driver for benthos diversity in these zones (Zettler et al., 2013). Despite the Benguela Upwelling System (BUS) being recognized as one of the major Eastern Boundary Upwelling systems, there is limited information on the structure and composition of the benthic communities. Information on macrofauna communities in the BUS will not only improve the existing database on benthic fauna but also provide insight into how increasing hypoxic areas in the ocean might structure benthic communities.

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In this study we adapted Levin's grouping system (Levin, 2003), classifying the different stations into three zones based on the oxygen levels recorded (microxic zone (<0.1 ml l⁻¹); dysoxic zone (0.1-1.0 ml 1⁻¹); oxic zone (>1.0 ml 1⁻¹)), which was supported by step-wise regression. The structure and composition of the macrofauna communities varied among the various oxygen zones. Most macrofaunal studies identify Polychaeta as the most abundant macrofauna taxon in both oxic and hypoxic areas (Eisenbarth & Zettler, 2016; Soto et al., 2017). However, in this study, Polychaeta only had the highest relative abundance in the oxic zones. In the microxic zone, the abundant taxon was Cumacea, followed by Polychaeta. It is essential to note that numerically Polychaeta was the most abundant in this oxygen zone, but the presence of other taxa in these stations reduced their relative abundance. . Here, relative abundance refers to the proportion of polychaetes to the total number of organisms in the same area. Therefore, even though polychaetes were numerically abundant, the diversity of other taxa present reduced their share of the total population, hence the low relative abundance. The presence of cumaceans in high abundance in the core OMZ (Area of lowest DO) has been reported by Zettler et al., (2013) and Eisenbarth & Zettler (2016), who described them as possible opportunistic species colonizing permanent hypoxic areas from adjacent areas, and thus their abundance may be season-specific. Currie et al. (2018) attributed the presence of Cumacea and other macrofauna taxa in the BUS to the Sulfuroxidizing bacteria, possibly providing a detoxified condition in this area. In this case, the mobility of the cumaceans gives them an advantage over other tolerant taxa like polychaetes and nematodes at the core and hence their high relative abundances at the OMZs core. Some polychaete families have physiological adaptations to tolerate the low oxygen quantities found in OMZs (Hanz et al., 2019; Joydas & Damodaran, 2014; Levin et al., 2009). At the microxic zone, polychaetes had the numerical abundance in the microxic stations despite cumaceans leading

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in relative abundance. In station (23002), where Cumacea were located, only two taxa were present. As a result, cumaceans accounted for 75% of the population despite having a density of only 932 individuals per square metre. In contrast, in station (23020), Polychaeta had the numerical high of 4350 individuals per square metre, which is relatively higher than the number recorded by cumaceans in station (23002). However, the diversity of other taxa in this station was much higher, and as a result, despite the high density, Polychaeta's proportion of the whole population was reduced to 53%, hence the lower relative abundance.

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In the dysoxic zone, the macro-nematoda was the most dominant taxa, outcompeting the polychaetes, in all the dysoxic stations, recording more than 70% relative abundance. Oxygen can cause shifts in community structure and trophic transfer (Neira et al., 2018), as evidenced in this study. Nematoda as a taxon has not received significant attention in the macrofaunal size range in most studies despite evidence of their presence therein (Joydas & Damodaran, 2014; Sharma et al., 2011). An increase in the size of nematodes to macrobenthic class sizes has been reported in chemosynthetic environments that experience similar characteristics as OMZs, i.e., low oxygen and high sulfidic contents (Vanreusel et al., 2010). Apart from the increase in size, OMZs also tend to enhance the regional dominance of tolerant organisms such as nematodes with high biomass recorded in response to organic matter inputs. The high abundances are thought to reflect the availability of organic matter, a significant nutrient source for macrofauna, coupled with a reduction in predation by, larger fauna that are affected adversely by the reduced oxygen concentrations (Moens et al., 2013).

The high nematode abundance in partnership with Oligochaeta occurred in the dysoxic zone; oxygen conditions were low enough to exclude some taxa but sufficient for tolerant species to survive and reproduce. Such conditions are referred to as the 'edge effect,' and such high densities

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579 are characteristics of the edge of the OMZs, where various species have been observed to have abnormally high densities. The reasons for these high abundances are not well understood, but 580 Gutiérrez et al. (2008) alluded that the nematodes' population can multiply in low oxygen 581 582 conditions, which experience high loads of organic matter input. Despite their tolerance to anoxia, 583 nematodes cannot survive long-term exposure, as observed in the anoxic zone (Moens et al., 2013). It is not clearly understood then whether the high abundance of the nematodes in one of the stations 584 is characteristic of the study site or just congregation to a food source. Thus, the patchiness and 585 high variability in the dysoxic zone calls for more studies (Buhl-Mortensen et al., 2010). 586 At dysoxic sites (DO 0.1-1.0 ml l⁻¹), other taxa like Ostracoda and Bivalvia were observed, albeit 587 in low numbers. Despite ostracodes flourishing better in well-oxygenated marine areas, various 588 589 families (Platycopina) have been observed to tolerate and thrive in Oxygen Minimum Zones. At oxic sites, where DO levels were above 1.0 ml l⁻¹, more taxa were recorded, and these numbers 590 increased even more when DO levels surpassed 2.0 ml 1⁻¹. 591 Our findings were in line with the various studies in OMZs whereby a decrease in richness and 592 diversity of macrofaunal species is observed towards the core and vice versa (McClain & 593 Schlacher, 2015). In the core (microxic) area, the macrofauna taxa had the Jowest density and 594 diversity. Each square meter of core area contained 1243 individuals, indicating low species 595 richness. A similar number was recorded by Zettler et al., (2009). In our study, we also observed 596 an increase in the number of taxa recorded in sites with DO levels above I ml l I The high 597 dominance of the tolerant taxa, as evidenced in the microxic and dysoxic areas, was reduced. Taxa 598 599 such as Amphipoda, Isopoda, Echinodermata, Nemertina, Aeolosomatidae, Aplacophora, Holothuroidea, and Cnidaria were only recorded in the oxic zones indicating low tolerance to low 600 601 oxygen levels. Of this fauna, crustaceans were the most abundant. This conforms to the

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et al. (2009) recorded amphipod species in low oxygen areas. These contradictory results indicate that, at least amongst the Amphipoda, tolerance/intolerance to hypoxia is species specific. Macro-nematoda abundance varied across the OMZ, with very low abundance in the microxic zones, extremely high numbers at the dysoxic zones, and a substantial amount at the oxic stations, accounting for only 28% of the total abundance. Nematodes are considered one of the most tolerant taxa in the marine environment, with the ability to tolerate low oxygen and high sulphidic environments characteristic of OMZs and may reach very high abundances in these environments (Neira et al., 2018; Gutierrez et al., 2008). This was the case at the dysoxic zone, where nematodes recorded high abnormal densities in one station, indicating the high tolerance of these taxa and their ability to dominate the macrofaunal component. Even with such high tolerance levels, nematode abundance can be impacted by microxic conditions, as observed in the microxic areas with a recording of only 621 nematodes m⁻² in this oxygen zone. This low, value, however, may be at the macrofauna level, and the case may be different at the meiofauna level, where nematodes have recorded substantial densities in microxic environments (Neira et al., 2018; Steyaert et al., 2007). Despite their high abundance in OMZs, not all nematodes are tolerant to low oxygen levels (Moens et al., 2013), as observed in this study. Metoncholaimus, Paracomesoma, and Desmolaimus dominated the dysoxic zone; these three genera are members of Oncholaimidae, Comesomatidae, and Linhomoeidae, respectively. Members of the family Oncholaimidae have large bodies that can disperse rapidly and colonize carcasses of macrofauna and even fish that may have succumbed to

the low levels of oxygen found in the dysoxic zone (Moens et al., 2013). Nevertheless, their bodies

are large enough to fit within the macrofauna size range, while their ability to swim ensures they

observations of Soto et al. (2017) at oxic sites in an upwelling system in Chile. Conversely, Zettler

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At the meiofaunal level (0.0038-1 mm), nematodes dominate with very high abundances in OMZs, and their structure and composition have been well documented (Gutiérrez et al., 2008; Neira et al., 2018). In contrast, despite various studies in OMZs acknowledging the presence of large nematodes (>1.00 mm), little attention has been accorded to analyzing

them further. In this study, we analyzed macro-Nematoda to

the genus level because of their dominance in the dysoxic zone to understand and acknowledge the structure and composition of macro-Nematoda in the BUS and OMZs in general.¶

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reflect a congregation upon a food source that had attracted nearby members in large numbers. Their ability to colonize the 'food source' in such numbers in a dysoxic environment indicates their tolerance to low oxygen levels.

On the other hand, Comesomatidae and Linhomoeidae members have been noted to have high abundances in enriched sediments with low oxygen, indicating tolerance to anoxic conditions (Steyaert et al., 2007). Their long and slender bodies might be the reason for their records at the macrofauna level. Despite this generalization at the family level and the assumptions that members of the same family may portray similar life strategies (Bongers et al., 1991), tolerance of nematodes to hypoxia is species-specific (Moens et al., 2013) as Steyaert et al. (2007) observed members of the same genera (Sabatiera) reacting differently to hypoxic and anoxic conditions. Thus, further

analysis should be done to identify the species that are tolerant to hypoxia at these OMZs at

macrofauna levels. Tolerance to hypoxia is indicated by both the presence and absence of

taxa most genera present in the oxic zone were absent in the dysoxic area and may be seen as

actively locate their food source (Moens et al., 2013). Their high abundance in this study might

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genera intolerant to low oxygen levels.

Wieser's feeding types, as outlined in his study (Wieser, 1953), have long been used to assess the trophic structure of nematode communities. This study had a high abundance of non-selective feeders (1B) and selective epigrowth feeders and herbivores (2A), with the latter dominating the dysoxic zone, while the former dominated the oxic zones.

Non-selective deposit feeders are commonly observed to dominate substrates characterized by high levels of organic matter and smaller grain size, indicating their adaptation to environments with elevated organic matter input, as observed in this study (Singh and Ingole, 2016; Heip et al.,

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714 1985). In contrast to previous findings on meiofaunal nematodes from Oxygen Minimum Zones 715 (OMZs) (Singh & Ingole, 2016; Neira et al., 2013) and macro-nematodes in deep-sea habitats (Sharma & Bluhm, 2011), which reported lower abundances of selective epigrowth feeders, our 716 717 study revealed a higher abundance of selective epigrowth feeders and herbivores. Within OMZs, such as our study site, the presence of giant bacteria such as Thioploca, Beggiatoa, and 718 719 Thiomargarita forming extensive mats (Brüchert et al., 2009) may provide a substantial food 720 source for nematodes (Giere, 2008). 721 Adjacent to OMZs, the prevalence of diatomaceous mud and the settling of fresh phytodetritus contribute significantly to the diet of epistratum feeders, leading to their high abundance in dysoxic 722 areas (Singh & Ingole, 2016; Sharma & Bluhm, 2011; Moens & Vincx, 1987; Heip et al., 1985). 723 724 However, in regions offshore from the OMZ, where the OMZ no longer interacts with the benthic zone, reduced diatom production diminishes the abundance of epistratum feeders, resulting in the 725 dominance of non-selective deposit feeders. 726 While non-selective deposit feeders typically dominate muddy and silty environments, selective 727 deposit feeders and carnivores have been observed to exhibit lower abundances in these habitats 728 (Heip et al., 1985). Our study aligns with this trend; however, other investigations have reported 729 730 contradictory findings, with selective deposit feeders being identified among the most dominant 731 nematodes in similar environments (Neira et al., 2013; Singh & Ingole, 2016). Additionally, carnivores and omnivores (2B)exhibited higher abundances in the dysoxic zone compared to the 732 oxic zone, with their overall abundance surpassing that of selective deposit feeders. These 733 observations suggest potential differences in nematode composition between the meiofaunal and 734 macrofaunal levels. Sharma and Bluhm (2011) observed that category 2B feeders were the second 735 736 most abundant feeding type in numerous stations within the Arctic deep-sea Canada Basin. They

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attributed these findings to the omission of larger nematodes in previous studies, as carnivores and

omnivores are often large-bodied species,

5.0 Conclusion

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In conclusion, this study provides insights into the intricate dynamics of macrofaunal communities in Oxygen Minimum Zones (OMZs). By classifying stations into microxic, dysoxic, and oxic zones based on oxygen levels, we observed variations in macrofaunal composition and abundance across these zones. Contrary to typical findings, Polychaeta dominated only in oxic zones, while Cumacea and Nematoda were prevalent in microxic and dysoxic zones, respectively. The presence of tolerant taxa like nematodes and Oligochaeta in dysoxic zones suggests their ability to thrive under low oxygen conditions. Conversely, taxa intolerant to low oxygen were absent in dysoxic zones, highlighting the importance of oxygen levels in shaping macrofaunal communities.

Macro-nematodes in general were the most abundant taxon. The high nematode densities suggest the ability of macro-nematodes to tolerate and thrive in low-oxygen environments. Our findings highlight the importance of considering macro-nematodes in studies of OMZ ecosystems and

underscore the need for further research to elucidate their ecological roles and responses to

environmental changes. Understanding the dynamics of macro-nematode populations coupled

with their meiofaunal co in OMZs is crucial for comprehensively assessing the impacts of oxygen

depletion on benthic communities and ecosystem functioning.

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Deleted: In summary, the Benguela Upwelling System (BUS) is a major Western Boundary Upwelling system known for its high surface productivity and Oxygen Minimum Zones (OMZ). This study found that the structure and composition of macrofauna communities in the BUS varied among different oxygen zones, with Polychaeta being the most abundant macrofauna taxon in anoxic zones, Cumacea dominating in microxic zones, Nematoda being the most dominant in dysoxic zones, and Ostracoda and Bivalvia observed limited numbers in the anoxic zone. These results suggest that oxygen levels significantly shape benthic communities, with tolerant taxaspecies dominating in lowoxygen environments and thus the increasing hypoxic areas in our ocean might result in reduced macrobenthic faunal densities, diversity, and species richness. Further studies are needed to understand the mechanisms behind the observed patchiness and high variability in the dysoxic zones.¶

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6.0 Tables

Table 1: Station information and abiotic factors information recorded from the stations in the Northern
 Benguela Upwelling System. N.R stands for no replicate.

Transect	Station (Replicate)	Longitude	Latitude	Depth (m)	том(%)	DO (ml l ⁻ ¹)
Off Terrace Bay						
(OTB-20°S)	20002(2)	12.99905	-20	33	27.47±1.55	1.8
	20020 <u>(3)</u>	12.67858	-20	125	38.64±2.16	0.67
	20040(N.R)	11.79321	-20	219	17.58 nr	0.53
Off Walvis Bay						
(OWB-23°S)	23002(NR)	14.3734	-23	39	23.08 nr	0.06
	23020(NR)	14.06986	-23	128	20.88 nr	0.02
	23070 <u>(3)</u>	13.14	-23	318	9.89±1.55	2.3
Off Luderitz (OL						
26°S)	26090(NR)	13.28	-26	1282	4.4 nr	4.34

Deleted: Cape Frio
Deleted: CF

Table 2: Sediment size distribution in the Northern Benguela Upwelling System

Transect	Station	Very Coarse Sand	Coarse Sand	Medium sand	fine sand	very fine sand	Silt
Off Terrace Bay (20°S)	20002	0.65	0.67	13.81	34.82	21.29	Deleted: Cape Frio
	20020	0.93	0.62	28.17	30.89	12.98	26.42
	20040	2.71	4.37	18.55	33.33	17.04	23.98
Off Walvis Bay (23°S)	23002	0.00	0.28	18.74	36.17	17.25	27.57
	23020	0.98	6.85	14.91	38.88	18.83	19.56
	23070	0.00	0.00	13.92	33.73	20.29	32.06
Off Luderitz (26 ^o S)	26090	0.00	0.29	23.99	30.46	19.25	26.01

7.0 Figures

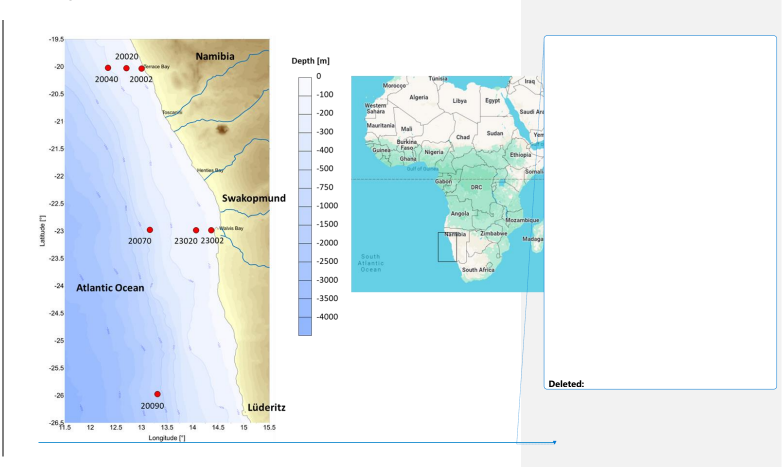


Figure 1: Sampling stations located across the Northern Benguela Upwelling System

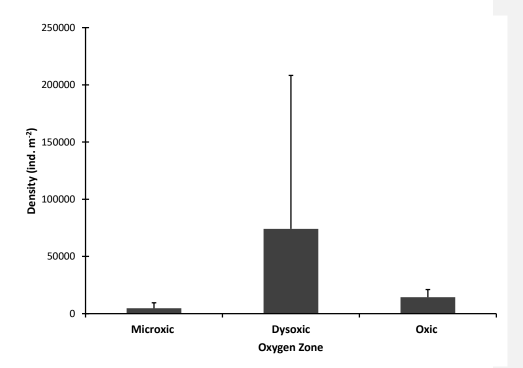


Figure 2: Macrobenthic densities in the different oxygen zones in the Northern Benguela Upwelling system

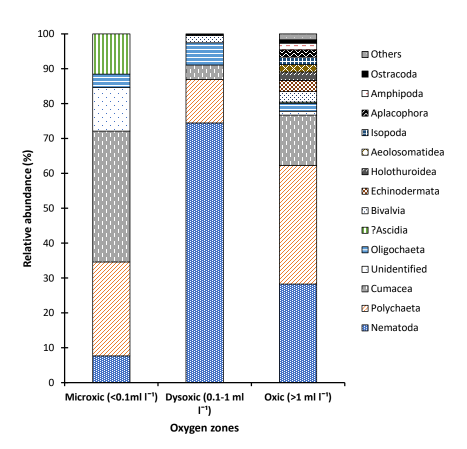


Figure 3: Macrofauna Relative abundance in the Northern Benguela Upwelling System, based on the oxygen zones identified.

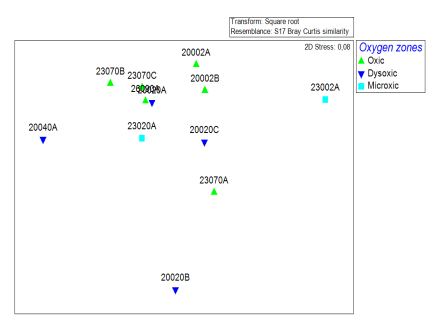


Figure 4: Non-metric multidimensional scaling (nMDS) plot based on Bray-Curtis Similarity index of macrobenthic fauna communities recorded in the Northern Benguela Upwelling System.

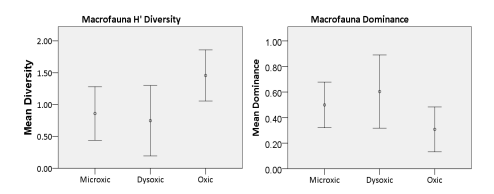


Figure 5: Macrofauna diversity indices recorded in the various oxygen zones in the Northern Benguela Upwelling system.

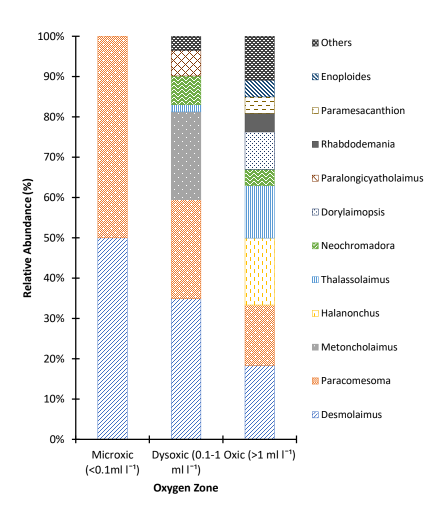


Figure 6: Macronematoda Relative abundance in the Northern Benguela Upwelling System, based on the oxygen zones identified.

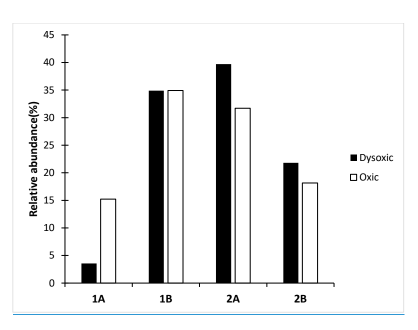


Figure 7: Feeding guilds based on Wieser of macro-Nematoda from the Northern Benguela Upwelling System. (1A= Selective deposit feeder, 1B- Non-Selective deposit feeder, 2A=Epigrwoth feeders, 2B=Predators/Omnivores).

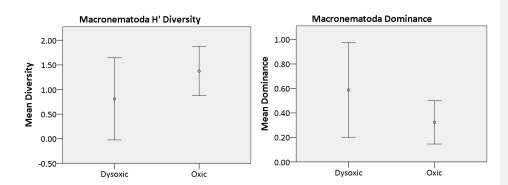


Figure 8: Macro-Nematoda diversity indices recorded in the various oxygen zones in the Northern Benguela Upwelling system.

8.0 Code/Data Availability

Currently, the data is not available but can be released upon request to the authors.

9.0 Author contribution

HS was responsible for the conceptualization of the study. HS and WB conducted the investigation, formal analysis, and drafting of the original draft. MA was responsible for supervision, review, and editing of the final draft.

10.0 Competing interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

11.0 Special Issue Statement

Part of this manuscript was presented as a poster presentation during the 53rd International Colloquium on Ocean Dynamics: 3rd GO2NE Oxygen Conference held in Liege, 2022. As a result, the corresponding author was invited to submit a manuscript for the special issue "Low-oxygen environments and deoxygenation in open and coastal marine waters" as part of the journal Biogeosciences.

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