# Long-term impacts of global temperature stabilization and overshoot on exploited marine species

Anne L. Morée<sup>1,2</sup>, Fabrice Lacroix<sup>1,2,3</sup>, William W. L. Cheung<sup>4</sup>, Thomas L. Frölicher<sup>1,2</sup>

<sup>1</sup>Climate and Environmental Physics, Physics Institute, University of Bern, Bern, Switzerland
 <sup>2</sup>Oeschger Centre for Climate Change Research, University of Bern, Bern, Switzerland
 <sup>3</sup>Institute of Geography, University of Bern, Bern, Switzerland
 <sup>4</sup>Institute for the Oceans and Fisheries, The University of British Columbia; Vancouver, Canada.

Correspondence to: Thomas L. Frölicher(thomas.froelicher@unibe.ch)

Abstract. Global warming alters ocean conditions, which can have dramatic consequences for marine species. Yet, the centennial-scale effects and reversibility of habitat viability for marine species, particularly those that are important to fisheries, remain uncertain. Using the Aerobic Growth Index, we quantify the impacts of warming and deoxygenation on the contemporary habitat volume of 46 exploited marine species in novel temperature stabilization and overshoot simulations until 2500. We demonstrate that only around half the simulated loss of contemporary (1995-2014) habitat volume is realized when warming levels are first reached. Moreover, in an overshoot scenario peaking at 2°C global warming before stabilizing at

15 1.5°C, the maximum decrease in contemporary habitat volume occurs more than 150 years post-peak warming. Species' adaptation may strongly mitigate impacts depending on adaptation rate and pressure. According to our study, marine species will be affected for centuries after temperature stabilization and overshoot, with impacts surpassing those during the transient warming phase.

#### 20 1 Introduction

Marine fisheries play a pivotal role in the global food system, generating economic benefits (Sumaila et al., 2012), supplying essential nutrients to humans (Hicks et al., 2019) and supporting people's livelihoods (Teh & Sumaila, 2013). Recent international assessments emphasize the predominately adverse impacts of climate change on marine ecosystems and fisheries (Bongaarts, 2019; IPCC, 2019, 2021). Projections indicate that ocean warming (Cheng et al., 2022) and deoxygenation
(Frölicher et al., 2009; Kwiatkowski et al., 2020; Schmidtko et al., 2017) will likely result in a global decline in total animal biomass (Lotze et al., 2019), potential fisheries catch (Cheung et al., 2016) and alterations in species distributions (Deutsch et al., 2020; Howard et al. 2020, Hodapp et al., 2023; Morée et al., 2023, Mongwe et al. 2024). Despite considerable research on the transient impacts of global warming and deoxygenation throughout the 21<sup>st</sup> century on marine fisheries, there remains a considerable gap in understanding the long-term repercussions of centennial-scale warming and deoxygenation. Moreover, the

30 increasing probability of a temporary temperature overshoot, despite efforts to limit global warming to the Paris Agreement's

1.5°C long-term goal (Rogelj et al., 2018), raises critical concerns - particularly regarding our limited understanding of the potential reversible or irreversible impacts of such a temporary overshoot on marine species (Meyer et al., 2022; Meyer & Trisos, 2023).

To comprehensively assess the multi-centennial impacts on marine species and potential reversibility after a temporary temperature overshoot, policy-relevant multi-centennial temperature stabilization and overshoot simulations are needed, but are currently lacking (King et al., 2021; "Reversing climate overshoot," 2023). Recent progress has been made in understanding the physical and biogeochemical reversibility and hysteresis under idealized scenarios (Frölicher & Joos, 2010; Jeltsch-Thömmes et al., 2020; Schwinger et al., 2022). These studies are, however, not extended to species impacts and are not designed to achieve specific global warming levels (Keller et al., 2018). Modelling studies that consider impacts on global

- 40 marine species are often limited to transient warming (i.e., a non-stabilized and therefore temporary warming and oxygen level (Cheung et al., 2016; Deutsch, Ferrel, Seible, et al., 2015; IPCC, 2019; Morée et al., 2023)) or idealized overshoot scenarios (Meyer et al., 2022; Santana-Falcón et al., 2023), and do not apply to specific species (Deutsch et al., 2020; Oschlies, 2021; Santana-Falcón et al., 2023). Moreover, modelling of species' distributions generally relies on the environmental state of the sea surface or sea floor only (Garciá Molinos et al., 2016; Heneghan et al., 2021; Hodapp et al., 2023; Meyer et al., 2022),
- 45 thereby neglecting the crucial vertical dimension when estimating changes in marine species distributions (GA, 2017) and hence largely neglecting changes in the ocean interior for disturbances in marine species distributions.

Recent advances in climate scenarios and impacts assessments on marine species under ocean warming and deoxygenation, such as the Adaptive Emission Reduction Approach (AERA; Terhaar et al., 2022) and the Aerobic Growth Index (AGI; Clarke et al., 2021; Morée et al., 2023), enable temperature-targeted modelling of the viability of contemporary distributions of marine

- 50 species. The AGI indicates the potential habitat conditions that theoretically support the aerobic scope required for the growth of marine water-breathing ectotherms, represented by the ratio of pO<sub>2</sub> supply to metabolic demand. AGI, when above the species-specific critical level (AGI<sup>crit</sup>), signifies the potential for a viable population of the considered species to be sustained with respect to temperature and the partial pressure of oceanic oxygen, pO<sub>2</sub>. This makes AGI a metric for habitat viability (Clarke et al., 2021; Morée et al., 2023). AERA achieves policy-relevant temperature stabilization at any desired level as well
- 55 as temporary temperature overshoot at any desired magnitude and duration by iteratively adjusting CO<sub>2</sub> forcing equivalent (CO<sub>2</sub>-fe) emissions in Earth system models. AGI, when above the species-specific critical level (AGI<sup>crit</sup>), signifies the potential for a viable population of the considered species to be sustained with respect to temperature and the partial pressure of oceanie oxygen, pO<sub>2</sub>. This makes AGI a metric for habitat viability (Clarke et al., 2021; Morée et al., 2023).

Here, we apply AERA to the Earth system model GFDL-ESM2M to simulate novel global warming scenarios spanning the

60 period 1861 to 2500, encompassing stable temperatures of 1.2°C, 1.5°C, 2°C, and 3°C as well as 2°C and 3°C overshoot scenarios followed by a return to stabilized 1.5°C warming relative to the 1861-1900 baseline (Fig. 1a,b). We quantify changes in the contemporary habitat volume of 46 representative marine species that are important to fisheries across the epipelagic,

mesopelagic, and demersal realms by utilizing AGI. Additionally, we examine the potential effects of species' adaptation to ocean warming and deoxygenation on their habitat volume loss, as adaptation is likely an important factor at the centennial

65 time scale considered here (Pinsky et al., 2020).

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#### 2 Methods

#### 2.1 Earth System Model

The temperature stabilization and overshoot simulations were carried out with the fully coupled Earth system model GFDL-ESM2M (Dunne et al., 2012, 2013). The GFDL-ESM2M couples an ocean, atmosphere, land, and sea-ice model. Ocean physics, including sea ice, is simulated with Modular Ocean Model version 4p1, at a 1° horizontal resolution which increases to up to <sup>1</sup>/<sub>3</sub>° meridionally at the equator and is tripolar above 65°N, with 50 vertical levels (Griffies, 2012). Mesoscale eddies are parameterized. Marine biogeochemistry is represented by Tracers of Ocean Productivity with Allometric Zooplankton version 2 (TOPAZv2) and consists of the cycles of carbon, nitrogen, phosphorus, silicon, iron, O<sub>2</sub>, alkalinity, and lithogenic particulate mattermaterial as well as surface sediment calcite. Three phytoplankton and one zooplankton group are explicitly

80 represented in TOPAZv2. Denitrification is included under suboxic conditions, while in the absence of both O<sub>2</sub> and nitrate additional respiration is accumulated as a negative O<sub>2</sub> concentration. The atmospheric module (AM2) consists of 24 vertical levels and a horizontal resolution of 2° latitude × 2.5° longitude (Anderson et al., 2004). The land model (LM3.0) represents the hydrological, energy and carbon cycles on land.

## 2.2 Model Simulations with the Adaptive Emissions Reduction Approach

- 85 The model employed the AERA (Terhaar et al., 2022) to conduct novel simulations spanning from 1861 to 2500 that stabilize global mean 2-m air temperature anomalies at 1.2°C, 1.5°C, 2°C, and 3°C, including temporary overshoots to 2°C and 3°C and then returning to stable 1.5°C warming relative to 1861-1900. The temperature changes were selected to align with the global warming levels commonly used in the IPCC reports 1.5 °C, 2.0 °C, and 3.0 °C (IPCC, 2022) as benchmarks for assessing impacts. Additionally, we included a stabilization scenario at the current level of warming to investigate committed impacts.
- 90 The warming rates leading to these temperature levels closely follow (at least initially) those observed during the historical period.

All simulations branch off an emissions-driven simulation over the historical period from 1861 to 2005. Post-2005, fossil fuel CO<sub>2</sub> emissions follow observed emissions until 2020 and projected emissions from the Nationally Determined Contributions

- 95 (Climate Action Tracker: https://climateactiontracker.org/global/temperatures/, last accessed December 2021) from 2021 to 2025. From 2025 onward, prescribed fossil-fuel CO<sub>2</sub> emissions are obtained every five years from AERA by subtracting CO<sub>2</sub> forcing equivalent (CO<sub>2</sub>-fe) emissions from prescribed non-CO<sub>2</sub> radiative agents and land use change from the AERA-derived total CO<sub>2</sub>-fe emission curve. Non-CO<sub>2</sub> radiative forcing and land use change are prescribed to follow the RCP2.6 scenario till 2100, with no changes thereafter. The AERA approach adapts CO<sub>2</sub>-fe emissions successively to converge to pre-defined
- 100 warming levels. AERA calculates every five years the realized simulated anthropogenic warming and the resulting remaining CO<sub>2</sub>-fe emission budget. This remaining emission budget is then distributed across future years using a cubic polynomial function. The temporary overshoot simulations are branched off from the 2°C and 3°C stabilization simulations, gradually reducing the warming level to 1.5°C by circa 2300 following the approach described in Terhaar et al. (2022). The resulting CO<sub>2</sub>-fe emission pathways that forced the model to the different warming levels show a maximum change in annual CO<sub>2</sub>-fe

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## maximum change in annual CO<sub>2</sub>-fe emissions of -0.7, and -1.6 Pg C yr<sup>-2</sup> for the 2°C and 3°C overshoot scenarios (Fig. A1).

emissions of -2.1, -1.5, -0.7, and -0.8 Pg C yr<sup>2</sup> for the 1.2°C, 1.5°C, 2°C, and 3°C stabilization scenarios respectively, and a

#### **2.3 Model Evaluation**

We briefly summarize previous evaluations of GFDL-ESM2M's physical and biogeochemical state with a focus on the variables most relevant for this study. In general, the model tends to better simulate physical than biogeochemical variables. The simulated ocean overturning structure and rate, ocean heat transport, ocean temperature and salinity, sea surface height, water mass age distributions and mixed layer depth show a general fair agreement with observational data (Dunne et al., 2012). Some potentially relevant mean biases include an overall too deep thermocline, Antarctic Bottom Water formation through open ocean convection, weak deep Pacific ventilation and too strong North Pacific thermocline ventilation (Dunne et al., 2012).

- 115 Furthermore, GFDL-ESM2M performs within the CMIP5 range with respect to O<sub>2</sub> projections, despite the global mean O<sub>2</sub> concentration being 5% lower than observed due to comparably large low-O<sub>2</sub> water volumes (Bopp et al., 2013). The spatial present-day distribution of O<sub>2</sub> agrees well with observations, and most of the bias in O<sub>2</sub> is limited to the Oxygen Minimum Zones in the eastern equatorial Pacific at depth (Frölicher et al., 2020), similar to other Earth system models (Cabré et al., 2015). The globally integrated NPP is 74 PgC yr<sup>-1</sup> in GFDL-ESM2M, compared to 53 Pg C yr<sup>-1</sup> on average in the observation-
- 120 based estimates (Le Grix et al. 2022). GFDL-ESM2M overestimates NPP, especially in the low latitudes, but despite these differences, GFDL-ESM2M succeeds in simulating the NPP mean spatial pattern of higher values in the low latitudes and lower values in the Southern Ocean and the subtropical gyres. As a result of the mean temperature and oxygen biases in the model, all mean biases have been corrected for by subtracting the mean bias between World Ocean Atlas 2018 data and the model from all model data (see below).

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A comparison of the relative changes in Aerobic Growth Index (AGI<sub>rel</sub>; see section 2.5) between CMIP6 model simulations and GFDL-ESM2M at transient 1.5°C warming allows additional evaluation of GFDL-ESM2M's performance. In CMIP6 models (Morée et al., 2023), the multi-model mean AGI<sub>rel</sub> changes are slightly smaller than in the GFDL-ESM2M, while patterns are generally similar. We attribute part of the relatively weaker signal in the CMIP6 ensemble compared to GFDL-

130 ESM2M to the later timing of reaching (transient) 1.5°C warming in the GFDL-ESM2M. Nevertheless, pelagic AGI<sub>rel</sub> largely falls within the CMIP6 range, while demersal AGI<sub>rel</sub> may be of larger magnitude than CMIP6.

#### 2.4 Model Data and Analysis

The annual mean GFDL-ESM2M data used in this study consist of dissolved O2 concentration, potential temperature, salinity,

- 135 ideal age, and the particulate organic carbon flux at 100m depth. The model data were first drift corrected by subtracting a linear trend obtained from the corresponding preindustrial control simulation over 1861-2500. The simulated O<sub>2</sub>, temperature and salinity data were bias-corrected using World Ocean Atlas observational data (WOA18; Garcia et al., 2019; Locarnini et al., 2018; Zweng et al., 2019) as follows: Climatological mean WOA18 data were regridded horizontally and vertically to the model grid after which the bias between the WOA18 data and the time mean model data over the years 1995-2014 were subtracted from the full time series. In-situ temperature is calculated from the potential temperature, and pO<sub>2</sub> is calculated from
- in-situ temperature and the salinity and  $O_2$  data (Bittig et al., 2018; Morée et al., 2023).

The 'first year of stable warming level' of its respective global warming level refers to the year in which the 31-yr running mean of the global mean 2-m air temperature is within one standard deviation of the aimed-for warming level (standard deviation of 0.074°C obtained from the preindustrial control simulation). The 31-yr mean was chosen to exclude interannual-to-decadal natural variability to a large degree. To make a fair comparison between the temperature stabilization scenarios with respect to the maximum amount of equilibration time available after reaching stable warming, committed habitat change is the change in the 296 years after hitting the respective stable warming level. Because Since the 3°C stabilization simulation reaches its warming level the latest <u>specifically in the year 2204 – there are 296 years remaining until the end of the simulation in 2500.</u>, it <u>Consequently, we adopt this determines this</u> 296-year timescale of for assessing committed changes in all

stabilization scenarios. Habitat change at a certain year is expressed as the forward 31-year mean at that year.

For the species-specific results, we estimate the combined uncertainty (e.g., whiskers in Fig. 3) from the timing of first reaching a stable warming level (varied by  $\pm 10$  years) and the species-specific control simulation variations in subcritical habitat

155 volume, which are combined using the Root Sum Square of the respective standard deviations. For the overshoot data, only the control standard deviation is considered as there is no stable warming level hit year.

#### 2.5 Aerobic Growth Index

We applied two alternative formulations of AGI as indicators of species-level and ecosystem level vulnerabilities to the impacts of ocean warming and deoxygenation (see eq. 1 and 2, respectively).

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For each species i, the Aerobic Growth Index (AGI) is calculated as the ratio between the pO<sub>2</sub> supply and demand (Eq. 1) (Clarke et al., 2021; Morée et al., 2023):

$$AGI_{i} = \frac{pO_{2}^{supply}}{pO_{2,i}^{demand}} = \frac{pO_{2}^{supply}}{pO_{2,i}^{threshold} \cdot (\frac{1}{3})^{1-d} \cdot exp\left(\frac{j_{2}-j_{1}}{r_{i}^{pref}}, \frac{j_{2}-j_{1}}{T}\right)},$$
(1)

1995-2014 time-mean AGI (Clarke et al., 2021; Morée et al., 2023).

with partial pressure  $pO_2$  (units in Pa) and in-situ temperature T (K). The generalized temperature dependence is represented by the variables  $j_1$  (the anabolism activation energy divided by the Boltzmann constant, 4500K) and  $j_2$  (the catabolism activation energy divided by the Boltzmann constant, 8000K), scaled by the metabolic scaling coefficient d (0.7). Aerobic

- activation energy divided by the Boltzmann constant, 8000K), scaled by the metabolic scaling coefficient d (0.7). Aerobic scope limits the species' distribution. Therefore, by using biogeographical data, we inferred the oxygen threshold necessary to support a viable population of the species. The species-specific critical threshold of  $pO_2$  ( $pO_{2,i}^{threshold}$ ) is calculated as the volume-weighted 10th percentile of 1995-2014 time-mean pO<sub>2</sub> within the 3-dimensional distribution of the species. The estimated pO<sub>2</sub> threshold for growth is proportional to critical pO<sub>2</sub> measured from physiological experiments (Clarke et al. 2021). The preferred in-situ temperature ( $T_i^{pref}$ ) is calculated as the volume-weighted 50th percentile of the 1995-2014 timemean temperature within the 3-dimensional distribution of the species, and  $AGI_i^{crit}$  as the volume-weighted 10th percentile of
- Using  $AGI_i$  and  $AGI_i^{crit}$ , we quantify the percentage of contemporary habitat available to sustain a viable population of the species (i.e., where  $AGI_i > AGI_i^{crit}$ ). Our use of this extended version of AGI (Morée et al., 2023) includes vertical variability in O<sub>2</sub> and temperature in both the estimate of species'  $pO_{2,i}^{threshold}$ ,  $T_i^{pref}$  and hence  $AGI_i$  and  $AGI_i^{crit}$ , as well as considering fully 3-dimensional changes in O<sub>2</sub> and temperature (as opposed to surface or sea floor values only). The generalized temperature dependence of  $pO_{2,i}^{demand}$  may cause over- or underestimation of an individual species' response to temperature.
- 180 The sensitivity of loss of contemporary habitat to this parameterization ( $j_2$ - $j_1$  in Eq. 1) is minor (Morée et al., 2023).

Changes in AGI between a time  $t_1$  and  $t_0$ , relative to  $t_0$  are species-independent and provide a sense of direction and magnitude of change in habitat viability (Morée et al., 2023):

$$AGI^{rel} = \frac{\Delta AGI}{AGI(t_0)} = \frac{pO_2(t_1)}{pO_2(t_0)} \cdot exp\left((j_2 - j_1) \cdot \left(\frac{1}{T(t_1)} - \frac{1}{T(t_0)}\right)\right) - 1$$
(2)

- 185 The t<sub>0</sub> refers to the mean over the period 1861-1900 in this study. Note that small changes in  $PO_2$  can lead to large changes in  $AGI_{rel}$  if reference  $PO_2$  (i.e.  $PO_2$  at t=0) is low (Eq. 2). Note that relative changes in AGI can be a poor indicator of the magnitude of change in contemporary habitat volume of specific species, for which species-specific critical thresholds are needed (Morée et al., 2023).
- 190 We recognize the potential habitat volume and the reversibility of their changes may not be realized by the species because of other biogeographic constraints such as dispersal potentials, availability of suitable prey or other environmental limitations beyond temperature and oxygen.

#### 2.6 Species data

Spatial distribution data for the 46 representative exploited species (species names are indicated in Figure 5; Morée et al., 2023; Palomares et al., 2004) are used to calculate  $T_i^{pref}$ ,  $pO_{2,i}^{threshold}$  and  $AGI_i^{crit}$ . These data also form the reference habitat 195 for assessing changes in contemporary habitat volume. The species were selected such that they provide a representative range in body size, climatic zone (tropical, temperate), habitat size, and depth range. The 46 species also cover a broad range of vulnerabilities to warming and deoxygenation, with the most vulnerable species having a  $\sim 30$  times larger change in volume per unit change in AGI than the least vulnerable species (Morée et al., 2023). We include 23 species with their predominant 200 occurrence in the epipelagic (0-200m depth), 5 species that mostly inhabit the mesopelagic (200-1000m depth) as well as 18 demersal species which live on or just above the sea floor (for which we use the deepest ocean model layer). Some pelagic and deep-water wide-ranging species were selected that inhabit both tropical and temperate regions. We assess contemporary habitat volume change by extending the 2D habitats over their depth range assuming that the distribution is the same throughout the water column. We acknowledge that some species may occupy only part of their assigned depth range or may temporarily 205 reside outside it, either above or below. Nevertheless, we believe that the assigned depth ranges generally provide a reasonable estimate of in-habitat  $pO_2$  and temperature variability. The reason for not just using the 3D model output across the entire depth range of each species' depth realm is primarily due to the current lack of reliable 3D species distributions for our selected species.

210 All habitat changes are changes in contemporary habitat volume since the 1861-1900 mean, to be consistent with the atmospheric temperature anomalies, as well as being presented as a percentage of viable contemporary habitat volume to facilitate comparison between species.

#### 2.7 Adaptation

Species will likely acclimatize and/or adapt to global warming by improving species' temperature and O<sub>2</sub> stress resistance on the timescales considered in this study (Pinsky et al., 2020). Such acclimatization and adaptation to environmental changes is already observed over the last few decades, for example for warm-water corals (Lachs et al., 2023; Logan et al., 2021). We base our adaptability assessment on a running climatology approach, which allows for the evaluation of both adaptation pressure to a species' habitat and the timescale of a species' adaptation to such pressure. We thereby follow the approach applied to corals by Logan et al., (2014), as outlined in the paragraphs below.<sup>27</sup>

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Adaptation pressure is evaluated by considering for each species a time-dependent  $AGI_{i,t}^{crit}$  ( $AGI_{i,t}^{crit}$ ) that changes when inhabitat AGI (and hence pO<sub>2</sub> and/or temperature) changes. As  $AGI_i$  and thus  $AGI_{i,t}^{crit}$  depends on  $T_i^{pref}$ ,  $pO_{2,i}^{threshold}$ , we also calculate a time-dependent  $pO_{2,i,t}^{threshold}$  ( $pO_{2,i,t}^{threshold}$ ) and  $T_i^{pref}$  ( $T_{i,t}^{pref}$ ). We do so by following the standard approach (described in section 2.5) but instead using a moving 20-year time-mean time window of pO<sub>2</sub>, temperature and AGI before each year t. In this way,  $AGI_{i,t}^{crit}$  is calculated for each species and each year over the period 1861 to 2500. It also follows that  $AGI_{i,t=2014}^{crit}$  is equal to the standard  $AGI_i^{crit}$ . For the first 20 years of our dataset, we use the time mean of all years available before and including that particular year.

We then consider the timescale of adaptation and its uncertainty by using mean  $AGI_{i,t}^{crit}$  over the 40, 60, 80, 100 (the range in 230 Logan et al., 2014), as well as 140, 180, and 220 years (to extend to species with longer adaptation times due to, for example, slower generational overturn) before year t in our quantification of contemporary habitat volume change at year t. A 40-year adaptation timescale thus means that a species has adjusted (i.e., adapted) its  $AGI_{i,t}^{crit}$  to the mean  $AGI_{i,t}^{crit}$  over the previous 40 years.

#### **3** Methods

#### 235 **3.1** Committed loss of contemporary habitat and overshoot impacts

In the stabilization simulations, the global atmospheric surface warming levels of 1.2°C, 1.5°C, 2°C, and 3°C are reached in the years 2019, 2054, 2091, and 2162 respectively (Fig. 1a). Afterwards, atmospheric temperatures are stabilized. Concurrently, the ocean is projected to warm and lose oxygen (Fig. 1c,e). By 2400-2500, the habitats of the 46 representative marine species have warmed by a median 0.8°C, 1.1°C, 1.5°C and 2.2°C (Fig. 1c), and lost 1.2 mbar, 1.3 mbar, 1.8 mbar, and

3.1 mbar of pO<sub>2</sub> in 2400-2500 for the 1.2°C, 1.5°C, 2°C, and 3°C stabilization scenarios, respectively (median across all species;
 Fig. 1e).



Figure 1: Simulated temperature stabilization and overshoot scenarios and associated impacts on median in-habitat temperature and pO<sub>2</sub> changes as well as on contemporary habitat volume across 46 representative marine species. (a,b) Global mean 2-m air temperature change since 1861-1900 for (a) temperature stabilization scenarios at 1.2°C, 1.5°C, 2°C, and 3°C warming, as well as (b) overshoot scenarios that peak at 2°C and 3°C warming before returning to 1.5°C warming. (c-h) Median change in (c,d) in-situ temperature and (e,f) pO<sub>2</sub> in the respective habitats of the 46 species and in (g,h) contemporary habitat volume (% of volume) since 1861-1900 for the (c,e,g) stabilization and (d,f,h) overshoot scenarios. Whiskers in (c-h)

250 indicate the 25<sup>th</sup> to 75<sup>th</sup> percentile range across the species habitat at the respective year of starting temperature stabilization and in year 2500. Thick lines show the median across the individual species' 31-year running mean data and thin lines the median across the species' their the annual mean time series.

With habitat warming and deoxygenation, species' habitat volume, defined here as the volume of contemporary habitat with the Aerobic Growth Index (AGI) above a species-specific threshold (see Methods), shrinks across the 46 species under all
stabilization scenarios (Fig. 1g). Contemporary habitat volume continues to decrease even after atmospheric temperatures have stabilized. As a result, only about half of the total loss of contemporary habitat volume is realized when first reaching the respective stable warming levels: The median fraction of realized habitat loss upon hitting stable warming levels, compared to the loss 296 years thereafter, amounts to 0.45 (with 25<sup>th</sup>-75<sup>th</sup> percentiles across the species at 0.19-0.76) for the 1.2°C scenario, 0.41 (0.25-0.67) for 1.5°C, 0.67 (0.36-0.93) for 2°C, and 0.60 (0.40-0.81) for the 3°C-stabilization scenario (Table 1). Adverse impacts on habitat viability of marine species thus continue to aggravate for centuries despite stabilization of the global mean atmospheric temperature.

Table 1. Simulated median habitat change (ΔHabitat) across the 46 representative species, presented in % of contemporary habitat volume at (a) time of warming level hit and (b) 296 years after time of warming level hit. (c)
Difference between (a) and (b), and (d) ratio between (a) and (b). In all columns the 25<sup>th</sup> and 75<sup>th</sup> percentile across the species is indicated in brackets and all values are 31-year forward running means.

Warming level	a. At time of warming	b. At time of warming	c. Committed (=b-a)	d. Ratio (=a/b)	
(°C)	level hit	level hit			
		+ 296 years			
1.2	-0.48 (-0.96, -0.34)	-1.06 (-2.32, -0.89)	-0.58 (-1.48, -0.28)	0.45 (0.19, 0.76)	
1.5	-0.76 (-1.32, -0.58)	-1.83 (-2.73, -1.31)	-1.07 (-1.72, -0.55)	0.41 (0.25, 0.67)	
2	-1.53 (-2.81, -1.15)	-2.28 (-3.92, -1.75)	-0.75 (-2.03, -0.06)	0.67 (0.36, 0.93)	
3	-2.78 (-4.40, -2.24)	-4.59 (-7.13, -3.14)	-1.82 (-2.94, -0.55)	0.60 (0.40, 0.81)	

∆Habitat (%)

After 296 years of temperature stabilization, the median decline in contemporary habitat among the species ranges from 1.06% (with 25<sup>th</sup>-75<sup>th</sup> percentiles across the species at 2.32% to 0.89%) for the 1.2°C stabilization scenario to 4.59% (7.13% to 3.14%) for the 3°C-stabilization scenario (Fig. 1g; Table 1). While these changes may appear small, the corresponding volumetric losses are substantial, ranging from thousands (order for the demersal species) to millions of km<sup>3</sup> (order for the epipelagic species), depending on the species and scenario (Morée et al., 2023). Notably, the 25% of species with the largest losses experience 2-5 times larger losses depending on the species and warming level (whiskers in Fig. 1g).

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Temporarily overshooting the 1.5°C warming level to 2°C or 3°C (Fig. 1b) drives bigger loss of contemporary habitat volume than direct stabilization at 1.5°C (Fig. 1h), due to associated overshoots of in-habitat warming (Fig. 1d) and deoxygenation (Fig. 1f). Peak loss of contemporary habitat volume during the 2°C and 3°C overshoot scenario is -2.1% and -3.2% in year 2270 and 2246, respectively (medians across the species, Table 2). This corresponds to -0.5% and -1.7% more loss of contemporary habitat than under the 1.5°C stabilization scenario at that time. Notably, peak loss of contemporary habitat is 280 realized 166 years (for the 2°C overshoot scenario) and 66 years (3°C overshoot) after maximum atmospheric warming in year 2104 and 2180 (Fig. 1b,h). By the time the maximum median loss of contemporary habitat is realized, atmospheric warming has therefore already returned to 1.7°C and 2.5°C, respectively. When atmospheric warming has returned to 1.5°C in 2400-2500, median contemporary habitats are 0.06% larger (0.31% smaller) in the 2°C (3°C) overshoot scenario than in 1.5°C 285 stabilization scenario. This indicates that for most species overshoot impacts are largely reversible by this time (Table 2). Nevertheless, the strong adverse impacts during the overshoot underscore the advantages of stabilizing at 1.5°C warming over overshooting this level.

Table 2. Simulated median habitat change ( $\Delta$ Habitat) across the 46 representative species, presented in % of contemporary habitat volume at (a) the year of maximum change for the respective scenario, (b) in 2400-2500, (c) the difference with the 1.5°C stabilization scenario in 2400-2500, and (d) the ratio relative to the 1.5°C stabilization

scenario in 2400-2500. In (d), the overshoot legacy is the difference between the overshoot scenario and the 1.5°C warming scenario relative to the change in the 1.5°C warming scenario. In all columns the 25<sup>th</sup> and 75<sup>th</sup> percentile across the species is indicated in brackets and all values are 31-year forward running means.

	AHabitat (%)				
Scenario	a. At peak change	b. In 2400-2500	c. Overshoot legacy in 2400-2500 minus stable 1.5°C	d. Overshoot legacy ratio in 2400-2500 relative to stable 1.5°C	
2°C overshoot	-2.13 (-3.11, -1.60)	-1.75 (-3.12, -1.24)	0.06 (-0.37, -0.03)	0.03 (-0.15, -0.03)	
3°C overshoot	-3.16 (-4.01, -2.42)	-2.12 (-3.66, -1.31)	-0.31 (-0.92, -0.10)	-0.17 (-0.36, -0.07)	

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#### 3.2 Spatial patterns and drivers of habitat viability changes

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To generalize these median impacts beyond the 46 analyzed species and to investigate the drivers of changes, we calculate the local relative changes in habitat viability expressed as the changes in the Aerobic Growth Index relative to the period 1861-1900 (AGI<sup>rel</sup>). These relative changes in habitat viability are fully species-independent and indicate the direction and extent of change in habitat viability across the world's oceans. We focus on the 1.5°C stabilization and the 2°C overshoot scenario, but results are qualitatively similar for the other scenarios.



Figure 2. Simulated committed change in the 1.5°C stabilization scenario (left panels) and overshoot legacy in the 2°C overshoot scenarios (right) in relative changes in habitat viability for three depth realms. (a-c) Committed change
in habitat viability is expressed as the difference between AGI<sup>rel</sup> at 2<u>9669</u> years after temperatures have stabilized and when the 1.5°C warming level is hit (i.e., the committed period). (d-f) Overshoot legacy in habitat viability is expressed as the

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difference between AGI<sup>rel</sup> in the 2°C overshoot and the 1.5°C stabilization scenarios in 2400-2500. Results are shown as the vertical mean over the epipelagic (0-200m) and mesopelagic (200-1000m) realms, and the demersal (sea floor) realm.

- 310 In the 1.5°C stabilization scenario, committed decreases in AGI<sup>rel</sup> (i.e., after temperatures have stabilized) intensify with depth and are comparable in magnitude to the changes during the period until 1.5°C warming is first reached (Fig. 2a-c). The most substantial committed declines in the epipelagic realm occur in the North Atlantic (Fig. 2a), driven by a sustained rise in ocean temperature (Fig. 3a) linked to the recovery of the Atlantic Meridional Overturning Circulation during the stabilization phase (Frölicher et al., 2020, Lacroix et al. 2024). Elsewhere in the pelagic and demersal realms, committed changes in habitat
- 315 viability are predominantly driven by O<sub>2</sub> changes through changes in apparent oxygen utilization (i.e., changes in ocean ventilation and organic matter remineralization) (Figs. 3 and A2). In the Pacific demersal realm (Fig. 2c), there's a committed reduction in habitat viability due to O<sub>2</sub> decrease tied to reduced deep ocean ventilation as indicated by older water ages (Fig. 3). Conversely, there is a committed increase in habitat viability in the North Pacific mesopelagic realm and Southern Ocean mesopelagic and demersal realms. The North Pacific increase is linked to increased ventilation and reduced biological oxygen
- 320 consumption (Figs. 2 and A2).-- The Southern Ocean increase is mainly driven by enhanced ventilation as indicated by younger water ages (Fig. 2).



Fig. 3. Simulated committed changes in the environmental state for the  $1.5^{\circ}$ C stabilization scenario for three depth realms. (a-c) Committed change in in-situ temperature, (d-f)  $pO_2$ , (g-i) AGI<sup>rel</sup> due to changes in O<sub>2</sub> only (calculated by keeping temperature constant at its 1861-1900 mean), (j-l) Apparent Oxygen Utilization (AOU= $O_2^{sat}-O_2$ ), and (m-o) ideal age of the water. As a proxy for change in ventilation time, we used the ideal age tracer, which is set to zero in the surface ocean and ages at a rate 1 yr yr<sup>-1</sup> below that. Committed change is the difference between environmental state at 296-269 years after temperatures have stabilized and when the 1.5°C warming level is hit. Results are shown as the vertical mean over the epipelagic (0-200m), mesopelagic (200-1000m), and demersal (sea floor) realms.

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The overshoot legacy for the 2°C overshoot is determined by comparing the changes in the overshoot scenario with the 1.5°C stabilization scenario without overshoot in the period 2400-2500, when global warming levels are similar (Fig. 2d-f). The most pronounced overshoot legacies are simulated in the northern flank of the eastern equatorial Pacific epipelagic (Fig. 2d) and in the mesopelagic of the North Pacific (Fig. 2e). These regions of habitat viability increase are caused by higher O<sub>2</sub> levels

335 resulting from reduced O<sub>2</sub> consumption and rejuvenating mesopelagic waters (Fig. 4j,k), which is consistent with previous idealized model simulations (Li et al., 2020; Santana-Falcón et al., 2023). In the demersal realm, relative changes in habitat viability are generally more negative under the overshoot than in the stabilization scenario by 2400-2500 (Fig. 2f). This negative legacy is mainly driven by reduced deep ocean ventilation as indicated by increased water ages (Fig. 4).



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**Fig. 4. Simulated overshoot legacy for the 2°C overshoot in the environmental state for three depth realms in 2400-2500.** (a-c) Overshoot legacy in in-situ temperature, (d-f)  $pO_2$ , (g-i) AGI<sup>rel</sup> due to changes in  $O_2$  only (calculated by keeping temperature constant at its 1861-1900 mean), (j-l) Apparent Oxygen Utilization (AOU= $O_2^{\text{sat}}$ - $O_2$ ), and (m-o) ideal age of the water. As a proxy for change in ventilation time, we used the ideal age tracer, which is set to zero in the surface ocean and ages at a rate 1 yr yr<sup>-1</sup> below that. Overshoot

345 legacy is the difference between state in the overshoot scenario and 1.5°C stabilization scenario in 2400-2500. Results are shown as the vertical mean over the epipelagic (0-200m), mesopelagic (200-1000m), and demersal (sea floor) realms.

#### 3.3 Impacts on contemporary habitat volume of individual species

- The committed loss of contemporary habitat volume is generally largest for demersal species (median of 1.7% in the 1.5°C stabilization scenario, and up to 6.4% for *Spectrunculus grandis*) and typically smallest in the epipelagic realm (median of 0.7%) (grey bars in Fig. 5a). When reaching stable 1.5°C warming in 2054, only a median 47% of the total loss is realized in the epipelagic, 56% in the mesopelagic and 28% in the demersal realm (Fig. A3a). Both the committed loss of contemporary habitat volume and the contrast in impact between the different depth realms generally strengthen with increased levels of global warming (Fig. A4). For most species, the committed loss of contemporary habitat volume is driven by ongoing decreases in dissolved O<sub>2</sub> (black stars in Fig. 5a). However, the magnitude of loss of contemporary habitat for individual species also
- depends on species-specific vulnerability and not only on the magnitude of changes in AGI<sup>rel</sup>, temperature or O<sub>2</sub> (Morée et al., 2023).



- 360 Figure 5. Simulated committed changes in contemporary habitat volume for the 1.5°C stabilization scenario (top panels) and overshoot legacy in the 2°C overshoot scenario (bottom) for 46 representative marine species and three different depth realms. (a) Committed change (grey bars) expressed as the difference between contemporary habitat volume changes at 29669 years after temperatures have stabilized and when the 1.5°C warming level is hit (i.e., the committed period). (b) Overshoot legacy (red bars) expressed as the difference between the 2°C overshoot scenario and the 1.5°C stabilization scenario in 2400-2500. Black stars in (a,b)
- 365 indicate contemporary habitat changes that are driven by temperature changes only (i.e., keeping O<sub>2</sub> values at 1861-1900 conditions). Maximum changes during the overshoot are indicated with red horizontal bars in (b). Whiskers in (a) indicate the uncertainty as the combined uncertainty coming from the uncertainty in timing of warming level hit and the species-specific control simulation variability (in (b), only the species-specific control simulation variability is considered in the whiskers as no stable warming level hit year is considered; see Methods).
- When temporarily overshooting stable 1.5°C warming to 2°C, median peak overshoot habitat loss exceeds the impact of 1.5°C warming by 1.1%, and at most 11.4% for *Thunnus atlanticus* (bars in Fig. 5b; see Fig. A4 for the 3°C overshoot). After overshoot returns in 2400-2500 from 2°C to stable 1.5°C warming, the species contemporary habitat volume is between -2.6% and +2.3% of stable 1.5°C warming (Fig. 5b), with a median loss of contemporary habitat volume of 0.10% in the epipelagic, 0.11% in the mesopelagic and 0.36% in the demersal realm (medians across depth realms in Fig. 5b). The overshoot legacy is
- 375 predominantly explained by enhanced (de)oxygenation in the overshoot simulation relative to the stabilization simulation in 2400-2500 (stars in Fig. 5b). Some species like *thunnus atlanticus* and *hippoglossus hippoglossus* even benefit from overshooting 1.5°C warming, mainly due to local oxygenation relative to the 1.5°C stabilization scenario.

#### 3.4 Potential adaptation

We consider both the adaptation pressure to a species which would incite adaptation (as the in-habitat change in habitat viability quantified by AGI) as well as the timescale of adaptation (the number of years needed to adapt a species' thresholds, evaluated for 40, 60, 80, 100, 140, 180, and 220 years). In consequence, after being under high (low) adaptation pressure a species will be relatively insensitive (sensitive) for a time that increases with the timescale of adaptation.





2°C warming level and in year 2500. Timeseries are smoothed with a 31-yr running mean.

Adaptation substantially reduces the projected loss of contemporary habitat compared to non-adaptation (Fig. 6). In the 2°Cstabilization scenario that includes adaptation, loss of contemporary habitat temporarily peaks at 0.63-1.53% around 2100 depending on the timescale of adaptation, before gradually recovering to 0.26-0.40% loss by 2500 (Fig. 6a, qualitatively similar results are obtained for the other scenarios). In comparison, the loss of contemporary habitat without adaptation is 2.54% by 2500. The maximum loss of contemporary habitat is larger and later for longer adaptation timescales. However, after the 21<sup>st</sup> century, loss of contemporary habitat in the adaptation scenarios is reduced again, because critical thresholds in AGI have had time to decrease enough (i.e., species could adapt to tolerate less favorable conditions).

In the overshoot scenarios, adaptation also causes reduced impact when compared to the non-adaptive scenarios (Fig. 6b). By 2400-2500, adaptation reduces the 3°C (2°C) overshoot legacy in terms of loss of contemporary habitat to 0.04% (0.28%), while this is 2.11% (1.75%) without adaptation. Following our approach, we find that the stronger the overshoot a species is exposed to, the lower the species' critical threshold becomes through adaptation (i.e., adapting to lower AGI<sup>crit</sup> and hence becoming more tolerant to low-oxygen or high-temperature environments). This results in a 'double overshoot' in the 3°C- overshoot simulation, where habitat volume impacts after ~2250 are smaller than in an adaptive 1.5°C stabilization scenario because species are still adapted to the (much) less favorable conditions in this recovery phase (Fig. 6b). In all scenarios with

adaptation, median impact on contemporary habitat volume does not reduce to zero because some species are exposed to a continued decrease in habitat viability even at the end of the 25<sup>th</sup> century.

#### **4 Discussion and Conclusions**

- 410 Through novel policy-relevant stabilization scenarios using an Earth system model, we show that only about half of the fish habitat changes have been realized when the temperature stabilization level is first hit. In overshoot scenarios, it may take over 150 years after the peak temperature for the most substantial impacts on marine species to manifest. Additionally, our findings suggest that rapid adaptation to changing conditions by species could mitigate some of the effects, contingent upon the rate of adaptation and external pressure.
- 415 An important assumption in our analysis is that the employed single Earth system model GFDL-ESM2M simulates changes in temperature and dissolved O<sub>2</sub> in a sufficiently realistic manner. We consider our global analysis for the warming and deoxygenation robust, especially given the good agreement of the model with observations and with other Earth system models (see Methods). However, projected local changes in oxygen may be less robust (Kwiatkowski et al., 2020), especially in low O<sub>2</sub> zones, and are likely to be underestimated (Buchanan & Tagliabue, 2021). Additional ESMs should be run under policy-
- 420 relevant temperature stabilization and overshoot scenarios to quantify the uncertainty in *p*O<sub>2</sub> changes. Nevertheless, multiple modelling studies indicate a centennial to multimillennial response timescale of O<sub>2</sub> to global warming (Battaglia & Joos, 2018; Bertini & Tjiputra, 2022; Frölicher et al., 2020), making it likely that multi-centennial O<sub>2</sub> changes are indeed important for committed impacts and overshoot legacies. Therefore, our foremost conclusion, that a major part of the total loss of contemporary habitat volume occurs after global temperatures have stabilized, likely does not depend on the model used but
- 425 the quantitative results may vary when using other models.

We highlight a multi-decadal to centennial delay in peak impact in contrast to peak global warming in the overshoot scenarios. This is much longer than the eight year delay found in Meyer et al., (2022) for marine species in a  $2^{\circ}$ C overshoot. Their use of sea surface data likely causes an underestimation of the peak delay due to faster reversibility of temperature and O<sub>2</sub> at the surface as compared to depth (Santana-Falcón et al., 2023; Schwinger et al., 2022), providing an additional argument for using

- 430 3-dimensional species distributions (GA, 2017). The depth of a species' distribution is also central to the quantification of the legacy impact of an overshoot after return to stable  $1.5^{\circ}$ C warming, since changes in ocean conditions in greater depths take substantially longer to reverse following overshoot scenarios reversibility timescales increase with depth (Lacroix et al. 2024, Santana-Falcón et al., 2023; Schwinger et al., 2022). Furthermore, the steep vertical gradients of  $pO_2$  and temperature as well as the dampened climate change signal at depth provide additional arguments to include the third, vertical, dimension when
- 435 estimating (changes in) contemporary species distributions (GA, 2017).

The Aerobic Growth Index faces limitations in its application due to several factors, notably the absence of species-specific fully three-dimensional distribution data, its neglect of critical stressors beyond warming and deoxygenation, and its confinement to assessing solely the loss of contemporary habitat. While we aim to capture habitat variability in  $O_2$  and temperature relevant for species-specific critical thresholds by extrapolating two-dimensional habitats across different depth

- 440 ranges, these approximations may not fully capture the complete picture. Despite this, our findings align with variable species responses seen in distribution shifts (Poloczanska et al., 2016), emphasizing the potential for substantial alterations in species compositions (Gotelli et al., 2022). Additionally, the response of marine organisms to global warming may encompass sensitivities to other effects, such as changes in low temperatures, acidity, nutrient availability, phenology, disease, predation pressure, invasive species, and (over)fishing (Gissi et al., 2021; Intergovernmental Panel on Climate Change (IPCC), 2023).
- 445 Multi-stressor research aims to assess and predict the cumulative effects of these stressors, including synergies and antagonistic relationships between them but is still bound by many challenges before full assessments can be made (Gissi et al., 2021; Orr et al., 2020). <u>T-he impacts of changing temperature and oxygen levels on species' habitat volume may be underestimated under the overshoot scenarios. Specifically, species may not be able to shift to viable habitats because of biogeographic constraints not represented by the AGI e.g., dispersal potential, trophic interactions. Such uncertainties are particularly notable for species</u>
- 450 with large transient changes in habitat volume such as blackfin tuna (*Thunnus atlanticus*) and those with relatively lower dispersal potential such as coral hind (*Cephalopholis miniata*). Furthermore, the AGI, in its current form, only represents the limitation of temperature and oxygen on the warm-temperature edge of fish distributions and does not represent the observed reduction in aerobic scope at the low-temperature edge (Pörtner 2010; Clarke et al. 2021; Deutsch et al. 2022). Thus, our results do not include habitat expansion or contraction due to ocean warming or cooling at the cold edge of species distributions.
- 455 Development and application of AGI that incorporate low temperature limitation of species' metabolism and growth will help quantify such uncertainties. Last, we note the importance of knowledge of species-specific critical  $pO_2$  thresholds and preferred temperature environments. Relative changes in  $pO_2$  supply over  $pO_2$  demand ratios have been implied to assess <u>ecosystemlevel</u> impact (Battaglia & Joos, 2018; Deutsch, Ferrel, Seibel, et al., 2015; Oschlies, 2021; Santana-Falcón et al., 2023), but species-specific thresholds and preference windows are needed for such estimates (Morée et al., 2023). Unlike the Metabolie

460 Index, which depends on laboratory estimates of critical thresholds, AGI is widely applicable to all species for which the distributions are known thanks to its generalized temperature dependence and species' distribution based critical thresholds. In the future, the application of the AGI to more species will allow the assessment of a wider range of interspecies responses.

Our sensitivity analysis shows that adaptation has large potential to mitigate the impacts of global warming on the habitat viability of marine fishes, although observations of a reduced equatorward extent of species' habitats suggest that adaptation

465 may be too slow, particularly for larger species such as marine fishes (Hastings et al., 2020). Furthermore, we strengthen the notion that the pathway as well as the magnitude of temperature overshoot is important for future ecological impact (Meyer & Trisos, 2023), and additionally show that warming pathways can further modulate impact through their effect on adaptation pressure. However, there is still a large gap in knowledge about marine fishes' adaptation to warming and deoxygenation.

Particularly, the rate of adaptation to warming and deoxygenation would depend on many biological and ecological factors

- that are not considered in this study e.g., the existing genetic diversity, species' life history traits, and population/meta-470 population structure and connectivity. In addition, one of many critical questions regarding how adaptation shapes biodiversity and biogeography under global change is the role of extreme events and tipping points in species adaptation. However, these dynamics remain challenging to study due to the scarcity of data on tipping events and evolutionary responses (Grant et al. 2017). Our analysis represents a first step in exploring the potential implications of adaptation for marine biogeography under
- centennial-scale climate change. Our findings could inspire future research to delve deeper into these complex and pressing 475 issues.

Commonly used CMIP6 overshoot scenarios projected a rapid rise to around 2°C of global warming, followed by reversal within 20-50 years (Pfleiderer et al., 2024). However, such a rapid reversal, requiring substantial atmospheric CO<sub>2</sub> removal, is unlikely given current limitations in carbon dioxide removal technologies (Schleussner et al., 2024). In our 3°C overshoot scenario, where we assume that CO<sub>2</sub> removal occurs over more than 150 to 200 years, negative CO<sub>2</sub>-fe emissions of up to 9 Pg C yr<sup>-1</sup> would still be required to bring temperatures back to 1.5°C (Fig. A1). This scale of removal far exceeds what is achievable in the foreseeable future (Fuss et al., 2018). Even in the 2°C overshoot scenario, the peak negative CO<sub>2</sub> emissions

would need to reach approximately 3 Pg C vr<sup>-1</sup>.

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We conclude that losses of marine species' contemporary habitats continue for centuries beyond reaching stable global warming levels and after peaking global warming in a temporary overshoot. Any impact assessed at transient warming levels (Hausfather et al., 2022; Morée et al., 2023) thus largely underestimates the total impact on marine species.

## **Appendix A: Additional figures**



490 Fig. A1. Prescribed annual CO<sub>2</sub> forcing equivalent emissions in the GFDL-ESM2M simulations. (a) CO<sub>2</sub>-fe emissions in the temperature stabilization scenarios. (b) CO<sub>2</sub>-fe emissions in the temperature overshoot scenarios and the reference 1.5°C stabilization scenario.



495 Fig. A2. Simulated committed changes in the 1.5°C stabilization (a) and overshoot legacy in the 2°C overshoot scenario (b) of particulate organic carbon export flux at 100m depth. (a) Committed changes are expressed as the ratio between the change during the committed period (the 29669 years after stable warming level hit) and at the time when the 1.5°C warming level is first hit. (b) Overshoot



legacy of the 2°C-overshoot scenario as the difference between the 2°C overshoot and the 1.5°C stabilization scenario divided by the 1.5°C stabilization scenario export flux in 2400-2500.

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Fig. A3. Simulated committed changes in the 1.5°C stabilization scenario (top panels) and overshoot legacy in the 2°C overshoot scenario (bottom panels) expressed in relative changes in contemporary habitat volume for 46 representative marine species and for three different depth realms. (a) Relative committed change as the ratio of habitat change at the time of stable warming level hit and the absolute total change in habitat after stabilization <u>296269</u> years later.

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Values of -100% or +100% (-100% is highlighted by a horizontal grey line) indicate that all change is realized at the time of

reaching the warming level. (b) Relative overshoot legacy (red bars) is expressed as the ratio between the 2°C overshoot scenario and the 1.5°C stabilization scenario in 2400-2500. *Hippoglossus hippoglossus* and *theragra chalcogramma* have a

relative overshoot legacy of 165% and 199%, respectively. Black stars in (a,b) indicate contemporary habitat changes that are driven by temperature changes only (i.e., keeping O<sub>2</sub> values at 1861-1900 conditions; see Methods). Whiskers in (a)

510 indicate the uncertainty as the combined uncertainty coming from the uncertainty in warming level hit timing and the

Epipelagic Demersal Mesopelagic а n **AHabitat** (%) -5 -10 1.2°C stabilization 2°C stabilization -15 3°C stabilization b 2°C overshoot 4 3°C overshoot ΔHabitat (%) 0 -2 -4 -6 \_۶ Aphanopus carbo Veocyttus rhomboidalis Makaira nigricans Brama brama Sarda sarda Acanthocybium solandri Elagatis bipinnulata Scomber japonicus atsuwonus pelamis Thunnus obesus stiophorus platypterus Thunnus alalunga Thunnus albacares Coryphaena hippurus Auxis thazard Alopias vulpinus Prionace glauca Alopias superciliosus Thunnus atlanticus Carcharhinus longimanus Euthynnus affinis Scomberomorus commerson Micromesistius poutassou Clupea harengus lhunnus maccoyii Sebastes mentella Hoplostethus atlanticus Spectrunculus grandis Carcharhinus falciformis Anarhichas denticulatus Cephalopholis miniata Decapterus russelli Beryx decadactylus Carangoides malabaricus Aelanogrammus aeglefinus Nezumia aequalis Epinephelus nigritus -epidocybium flavobrunneum Bothus pantherinus Lutjanus argentimaculatus Hippoglossus hippoglossus Hippoglossoides platessoides Centrolophus nige Beryx splenden Gadus mor Theragra chalcogram

species-specific control simulation variability (in (b), only the species-specific control simulation variability is considered in the whiskers as no warming level hit year is considered; see Methods).

- 515 Fig. A4. Simulated changes in contemporary habitat volume (%) for 46 representative marine species for different stabilization (top panels) and overshoot (bottom) scenarios. (a) Committed change expressed as the difference between contemporary habitat volume changes at 29669 years after temperatures have stabilized and when the 1.5°C warming level is hit (i.e., the committed period). (b) Overshoot legacy as the difference between the 2°C or 3°C overshoot and 1.5°C stabilization scenario in 2400-2500. Maximum changes during the overshoot (10-year running mean filter) are indicated with beriarneta have. Whiskers in (a) indicate the uncertainty as the combined uncertainty coming from the uncertainty in the uncertainty is the combined uncertainty coming from the uncertainty in the uncertainty in the uncertainty is the combined uncertainty coming from the uncertainty in the uncertainty is the combined uncertainty is the uncertainty in the uncertainty is the combined uncertainty is the uncertainty in the uncertainty is the combined uncertainty is the uncertainty in the uncertainty is the combined uncertainty is the uncertainty is the combined uncertainty is the uncertainty in the uncertainty is the uncertainty is the combined uncertainty is the u
- 520 horizontal bars. Whiskers in (a) indicate the uncertainty as the combined uncertainty coming from the uncertainty in warming level hit timing and the species-specific control simulation variability (in (b), only the species-specific control simulation variability is considered in the whiskers as no warming level hit year is considered; see Methods).

## 525

## Code and Data Availability

Data and scripts for reproduction of the manuscript figures are freely accessible at https://doi.org/10.5281/zenodo.14504902.

## Author contribution

530 AM and TF designed the study. FL conducted the Earth system simulations. WWLC was instrumental in developing the Aerobic Growth Index. AM performed the analysis and wrote the first draft. All authors contributed to the writing of the manuscript.

#### **Competing interests**

535 The authors have not conflicts of interest to declare.

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#### 545 **References**

Anderson, J. L., Balaji, V., Broccoli, A. J., Cooke, W. F., Delworth, T. L., Dixon, K. W., Donner, L. J., Dunne, K. A., Freidenreich, S. M., Garner, T., Gudgel, R. G., Gordon, C. T., Held, I. M., Hemler, R. S., Horowitz, L. W., Klein, S. A., Knutson, T. R., Kushner, P. J., Langenhorst, A. , R., Lau, N.-C., Liang, Z., Malyshev, S. L., Milly, P. C. D., Nath, M. J., Ramaswamy, V., Schwarzkopf, M. D., Shevliakova, E., Sirutis, J. J., Soden, B. J., Stern, W. F., Thompson, L. A., Wilson, R. J., Wittenberg, A. T., Wyman, B. L.: The new GFDL global atmosphere and land model AM2-LM2: Evaluation with prescribed

550 J., Wittenberg, A. T., Wyman, B. L.: The new GFDL global atmosphere and land model AM2-LN SST simulations. J. Climate, 17(24), 4641–4673. https://doi.org/10.1175/JCLI-3223.1, 2004.

Battaglia, G., & Joos, F.: Hazards of decreasing marine oxygen: the near-term and millennial-scale benefits of meeting the Paris climate targets. Earth Syst. Dynam., 9(2), 797–816. <u>https://doi.org/10.5194/esd-9-797-2018</u>, 2018.

Bertini, L., & Tjiputra, J.: Biogeochemical Timescales of Climate Change Onset and Recovery in the North Atlantic Interior 555 Under Rapid Atmospheric CO<sub>2</sub> Forcing. J. Geophys. Res.-Oceans, 127(4). <u>https://doi.org/10.1029/2021JC017929</u>, 2022.

Bittig, H., Körtzinger, A., Johnson, K., Claustre, H., Emerson, S., Fennel, K., Garcia, H., Gilbert, D., Gruber, N., Kang, D.-J., Naqvi, W., Prakash, S., Riser, S., Thierry, V., Tilbrook, B., Uchida, H., Ulloa, O., Xing, X.: SCOR WG 142: Quality Control Procedures for Oxygen and Other Biogeochemical Sensors on Floats and Gliders. Recommendations on the conversion

between oxygen quantities for Bio-Argo floats and other autonomous sensor platforms. CNRS, UMR 7093, Laboratoire d'Océanographie de Villefranche (LOV), Villefranche sur Mer, France: Ifremer. https://doi.org/10.13155/45915, 2018.

Bongaarts, J.: IPBES, 2019. Summary for policymakers of the global assessment report on biodiversity and ecosystem services of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services. Population and Development Review, 45(3). <u>https://doi.org/10.1111/padr.12283</u>, 2019.

Bopp, L., Resplandy, L., Orr, J. C., Doney, S. C., Dunne, J. P., Gehlen, M., Halloran, P., Heinze, C., Ilyina, T., Séférian, R.,
Tjiputra, J., Vichi, M.: Multiple stressors of ocean ecosystems in the 21st century: Projections with CMIP5 models. Biogeosciences, 10(10), 6225–6245. https://doi.org/10.5194/bg-10-6225-2013, 2013.

Buchanan, P. J., & Tagliabue, A.: The Regional Importance of Oxygen Demand and Supply for Historical Ocean Oxygen Trends. Geophys. Res. Lett., 48(20). <u>https://doi.org/10.1029/2021GL094797</u>, 2021.

Cabré, A., Marinov, I., Bernardello, R., & Bianchi, D.: Oxygen minimum zones in the tropical Pacific across CMIP5 models:
 Mean state differences and climate change trends. Biogeosciences, 12(18), 5429–5454. <u>https://doi.org/10.5194/bg-12-5429-2015</u>, 2015.

Cheng, L., von Schuckmann, K., Abraham, J. P., Trenberth, K. E., Mann, M. E., Zanna, L., England, M., H., Zika, J. D., Fasullo, J. T., Yu, Y., Pan, Y., Zhu, J., Newsom, E. R., Bronselaer, B., Lin, X.: Past and future ocean warming. Nature Reviews Earth and Environment. <u>https://doi.org/10.1038/s43017-022-00345-1</u>, 2022.

575 Cheung, W. W. L., Reygondeau, G., & Frölicher, T. L.: Large benefits to marine fisheries of meeting the 1.5°C global warming target. Science, 354(6319), 1591–1594. <u>https://doi.org/10.1126/science.aag2331</u>, 2016.

Clarke, T. M., Wabnitz, C. C. C., Striegel, S., Frölicher, T. L., Reygondeau, G., & Cheung, W. W. L.: Aerobic growth index (AGI): An index to understand the impacts of ocean warming and deoxygenation on global marine fisheries resources. Progress in Oceanography, 195, 102588. <u>https://doi.org/10.1016/j.pocean.2021.102588</u>, 2021.

580 Deutsch, C., Ferrel, A., Seibel, B., Pörtner, H.-O., & Huey, R. B.: Climate change tightens a metabolic constraint on marine habitats. Science, 348(6239), 1132–1135. https://doi.org/10.1126/science.aaa1605, 2015.

Deutsch, C., Penn, J. L., & Seibel, B.: Metabolic trait diversity shapes marine biogeography. Nature, 585(7826). https://doi.org/10.1038/s41586-020-2721-y, 2020.

Dunne, J. P., John, J. G., Adcroft, A. J., Griffies, S. M., Hallberg, R. W., Shevliakova, E., Stouffer, R. J., Cooke, W., Dunne,
K. A., Harrison, M. J., Krasting, J. P., Malyshev, S. L., Milly, P. C. D., Phillipps, P. J., Sentman, L. T., Samuels, B. L., Spelman,
M. J., Winton, M., Witternberg, A. T., Zadeh, N.: GFDL's ESM2 Global Coupled Climate-Carbon Earth System Models. Part

I: Physical Formulation and Baseline Simulation Characteristics. J. Climate, 25, 6646–6665. <u>https://doi.org/10.1175/JCLI-D-11-00560.1</u>, 2012.

Dunne, J. P., John, J. G., Shevliakova, S., Stouffer, R. J., Krasting, J. P., Malyshev, S. L., Milly, P. C. D., Sentman, L. T.,
Adcroft, A. J., Cooke. W., Dunne, K. A., Griffies, S. M., Hallberg, R. W., Harrison, M. J., Levy, H., Wittenberg, A. T., Phillips,
P. J., Zadeh, N. GFDL's ESM2 global coupled climate-carbon earth system models. Part II: Carbon system formulation and baseline simulation characteristics. J. Climate, 26(7), 2247–2267. https://doi.org/10.1175/JCLI-D-12-00150.1, 2013.

Frölicher, T. L., Joos, F., Plattner, G. K., Steinacher, M., & Doney, S. C.: Natural variability and anthropogenic trends in oceanic oxygen in a coupled carbon cycle-climate model ensemble. Global Biogeochemical Cycles, 23(GB1003), 1–15.
 https://doi.org/10.1029/2008GB003316, 2009.

Frölicher, T L, Aschwanden, M. T., Gruber, N., Jaccard, S. L., Dunne, J. P., & Paynter, D.: Contrasting Upper and Deep Ocean Oxygen Response to Protracted Global Warming. Global Biogeochemical Cycles, 34(8), e2020GB006601. https://doi.org/https://doi.org/10.1029/2020GB006601, 2020.

Frölicher, Thomas L., & Joos, F.: Reversible and irreversible impacts of greenhouse gas emissions in multi-century projections with the NCAR global coupled carbon cycle-climate model. Climate Dynamics, 35(7), 1439–1459. https://doi.org/10.1007/s00382-009-0727-0, 2010.

Fuss, S., Lamb, W. F., Callaghan, M. W., Hilaire, J., Creutzig, F., Amann, T., Beringer, T., de Oliveira Garcia, W., Hartmann, J., Khanna, T.: Negative emissions - Part 2: Costs, potential and side effects. Environmental Research Letters, 13, 063002. https://doi.org/10.1088/1748-9326/aabf9f, 2018.

605 Duffy, G. A., Chown, S. L.: Explicitly integrating a third dimension in marine species distribution modelling. Marine Ecology Progress Series, 564, 1–8. Retrieved from <u>https://www.int-res.com/abstracts/meps/v564/p1-8/</u>, 2017.

Grant, P. R., Grant, B. R., Huey, R. B., Johnson, M. T. J., Knoll, A. H., Schmitt, J.: Evolution caused by extreme events. Philosophical Transactions of the Royal Society B: Biological Sciences, 372(1723), 20160146. https://doi.org/10.1098/rstb.2016.0146, 2017.

610 Garcia, H. E., Weathers, K. W., Paver, C. R., Smolyar, I., Boyer, T. P., Locarini, R. A., Zweng, M. M., Mishonov, A. B., Baranova, O. K., Seidov, D., Reagan, J. R.: World Ocean Atlas 2018, Volume 3: Dissolved Oxygen, Apparent Oxygen Utilization, and Dissolved Oxygen Saturation. World Ocean Atlas 2018, 83(July), 38pp. 2019.

615

Garciá Molinos, J., Halpern, B. S., Schoeman, D. S., Brown, C. J., Kiessling, W., Moore, P. J., Pandolfi, J. M., Poloczanska, E. S., Richardson, A. J., Burrows, M. T., Climate velocity and the future global redistribution of marine biodiversity. Nature Climate Change, 6(1). https://doi.org/10.1038/nclimate2769, 2016.

Gissi, E., Manea, E., Mazaris, A. D., Fraschetti, S., Almpanidou, V., Bevilacqua, S., Coll, M., Guarnieri, G., Loret-Lloret, E., Pascual, M., Petza, D., Rilov, G., Schonwald, M., Stelzenmüller, V., Katsanevakis, S.: A review of the combined effects of

climate change and other local human stressors on the marine environment. Science of the Total Environment. https://doi.org/10.1016/j.scitotenv.2020.142564, 2021.

620 Gotelli, N. J., Moyes, F., Antão, L. H., Blowes, S. A., Dornelas, M., McGill, B. J., Penny, A., Schipper, A. M., Shimadzu, H., Supp, S. R., Waldock, C. A., Magurran, A. E.: Long-term changes in temperate marine fish assemblages are driven by a small subset of species. Global Change Biology, 28(1), 46–53. <u>https://doi.org/https://doi.org/10.1111/gcb.15947</u>, 2022.

Griffies, S. M.: Elements of the Modular Ocean Model (MOM). GFDL Model Documentation, 3(C), 1-631, 2012.

Hastings, R. A., Rutterford, L. A., Freer, J. J., Collins, R. A., Simpson, S. D., & Genner, M. J., 2020, Climate Change Drives
Poleward Increases and Equatorward Declines in Marine Species. Current Biology, 30(8). https://doi.org/10.1016/j.cub.2020.02.043, 2020.

Hausfather, Z., Marvel, K., Schmidt, G. A., Nielsen-Gammon, J. W., & Zelinka, M.: Climate simulations: recognize the 'hot model' problem. Nature. <u>https://doi.org/10.1038/d41586-022-01192-2</u>, 2022.

- Heneghan, R. F., Galbraith, E., Blanchard, J. L., Harrison, C., Barrier, N., Bulman, C., Cheung, W., Coll, M., Eddy, T. D.,
- 630 Erauskin-Extramiana, M., Everett, J. D., Fernades-Salvador, J. A., Gascuel, D., Guiet, J., Maury, O., Palacios-Abrantes, J., Petrik, C. M., du Pontavice, H., Richardson, A. J., Steenbeek, J., Tai, T. C., Volkholz, J., Woodworth-Jefcoats, P. A., Tittensor,

D. P.: Disentangling diverse responses to climate change among global marine ecosystem models. Progress in Oceanography, 198. <u>https://doi.org/10.1016/j.pocean.2021.102659</u>, 2021.

Hicks, C. C., Cohen, P. J., Graham, N. A. J., Nash, K. L., Allison, E. H., D'Lima, C., Mills, D. J., Roscher, M., Thilsted, S.
H., Thorne-Lyman, A. L., MacNeil, M. A.: Harnessing global fisheries to tackle micronutrient deficiencies. Nature, 574(7776), 95–98. https://doi.org/10.1038/s41586-019-1592-6, 2019.

Hodapp, D., Roca, I. T., Fiorentino, D., Garilao, C., Kaschner, K., Kesner-Reyes, K., Schider, B., Segschneider, J., Kocsis, A. T., Kiessling, W., Brey, T., Froese, R.: Climate change disrupts core habitats of marine species. Global Change Biology, 29(12). <u>https://doi.org/10.1111/gcb.16612</u>, 2023.

640 Howard, E. M., Penn, J. L., Frenzel, H., Seibel, B. A., Bianchi, D., Renault, L., Kessouri, F., Sutula, M. A., Mcwilliams, J. C., and Deutsch, C.: Climate-driven aerobic habitat loss in the California Current System, Sci. Adv., 6, eaay3188, https://doi.org/10.1126/sciadv.aay3188, 2020.

Intergovernmental Panel on Climate Change (IPCC): Oceans and Coastal Ecosystems and Their Services. In Climate Change 2022 – Impacts, Adaptation and Vulnerability. <u>https://doi.org/10.1017/9781009325844.005</u>, 2023.

645 IPCC: The Ocean and Cryosphere in a Changing Climate. Intergovernmental Panel on Climate Change, (September), 1–765. https://doi.org/https://www.ipcc.ch/report/srocc/, 2019.

IPCC: Summary for Policymakers. In V. Masson-Delmotte, P. Zhai, A. Pirani, S. L. Connors, C. Péan, S. Berger, et al. (Eds.), Climate Change 2021: The Physical Science Basis. Contribution of Working Group I to the Sixth Assessment Report of the

Intergovernmental Panel on Climate Change (p. 3–32). Cambridge, United Kingdom and New York, NY, USA: Cambridge University Press. https://doi.org/10.1017/9781009157896.001, 2021.

IPCC: IPCC, 2022: Summary for Policy Makers. Climate Change 2022: Impacts, Adaptation and Vulnerability, 2022.

Jeltsch-Thömmes, A., Stocker, T. F., & Joos, F.: Hysteresis of the Earth system under positive and negative CO2 emissions. Environmental Research Letters, 15(12). <u>https://doi.org/10.1088/1748-9326/abc4af</u>, 2020.

Keller, D. P., Lenton, A., Scott, V., Vaughan, N. E., Bauer, N., Ji, D., Jones, C. D., Kravitz, B., Muri, H., Zickfeld, K.: The
 Carbon Dioxide Removal Model Intercomparison Project (CDRMIP): Rationale and experimental protocol for CMIP6.
 Geoscientific Model Development, 11(3), 1133–1160. <u>https://doi.org/10.5194/gmd-11-1133-2018</u>, 2018.

King, A. D., Sniderman, J. M. K., Dittus, A. J., Brown, J. R., Hawkins, E., & Ziehn, T.: Studying climate stabilization at Paris Agreement levels. Nature Climate Change. <u>https://doi.org/10.1038/s41558-021-01225-0</u>, 2021.

Kwiatkowski, L., Torres, O., Bopp, L., Aumont, O., Chamberlain, M., Christian, J. R., Dunne, J., P., Gehlen, M., Ilyina, T.,

660 John, J. G., Lenton, A., Li, H., Lovenduski, N. S., Orr, J. C., Palmieri, J., Santana-Falcon, Y., Schwinger, J., Séférian, R., Stock, C. A., Tagliabue, A., Takano, Y., Tjiputra, J., Toyama, K., Tsujino, H., Watanabe, M., Yamamoto, A., Yool, A., Ziehn, T.: Twenty-first century ocean warming, acidification, deoxygenation, and upper-ocean nutrient and primary production decline from CMIP6 model projections. Biogeosciences, 17(13), 3439–3470. <u>https://doi.org/10.5194/bg-17-3439-2020</u>, 2020.

Lachs, L., Donner, S. D., Mumby, P. J., Bythell, J. C., Humanes, A., East, H. K., & Guest, J. R.: Emergent increase in coral
 thermal tolerance reduces mass bleaching under climate change. Nature Communications, 14(1).
 <a href="https://doi.org/10.1038/s41467-023-40601-6">https://doi.org/10.1038/s41467-023-40601-6</a>, 2023.

Lacorix, F., Burger, F. A., Silvy, Y., Schleussner, C.-F., Frölicher, T. L.: Persistently elevated high-latitude ocean temperatures and global sea level following temporary temperature overshoots. Earth's Future, 12, e2024EF004862. https://doi.org10.1029/2024EF004862, 2024.

670 Li, X., Zickfeld, K., Mathesius, S., Kohfeld, K., Matthews, J. B. R.: Irreversibility of Marine Climate Change Impacts Under Carbon Dioxide Removal. Geophysical Research Letters, 47(17). <u>https://doi.org/10.1029/2020GL088507</u>, 2020.

Locarnini, R. A., Mishonov, A. V., Baranova, O. K., Boyer, T. P., Zweng, M. M., Garcia, H. E., Reagan, J. R., Seidov, D., Weathers, K. W., Paver, C. R., Smolyar, I. V.: World Ocean Atlas 2018, Volume 1: Temperature. NOAA Atlas NESDIS 81, 1(July), 2019.

675 Logan, C. A., Dunne, J. P., Eakin, C. M., & Donner, S. D.: Incorporating adaptive responses into future projections of coral bleaching. Global Change Biology, 20(1), 125–139. <u>https://doi.org/10.1111/gcb.12390</u>, 2014.

Logan, C. A., Dunne, J. P., Ryan, J. S., Baskett, M. L., & Donner, S. D.: Quantifying global potential for coral evolutionary response to climate change. Nature Climate Change, 11(6). <u>https://doi.org/10.1038/s41558-021-01037-2</u>, 2021.

Lotze, H. K., Tittensor, D. P., Bryndum-Buchholz, A., Eddy, T. D., Cheung, W. W. L., Galbraith, E. D., Barange, M., Barrier,
N., Bianchi, D., Blanchard, J. L., Bopp, L., Büchner, M., Bulman, C. M., Carozza, D. A., Christensen, V., Coll, M., Dunne, J.
P., Fulton, E. A., Jennings, S., Jones, M. C., Mackinson, S., Maury, O., Niiranen, S., Oliveros-Ramos, R., Roy, T., Fernades,
J. A., Schewe, J., Shin, Y.-J., Silva, T. A. M., Steenbeek, J., Stock, C. A., Verley, P., Volkholz, J., Walker, N., D., Worm, B.:

Global ensemble projections reveal trophic amplification of ocean biomass declines with climate change. Proceedings of the National Academy of Sciences, 116(26), 12907 LP – 12912. <u>https://doi.org/10.1073/pnas.1900194116</u>, 2019.

685 Meyer, A. L. S., & Trisos, C. H.: Ecological impacts of temperature overshoot: The journey and the destination. One Earth, 6(12), 1614–1617. <u>https://doi.org/10.1016/j.oneear.2023.11.014</u>, 2023.

Meyer, A. L. S., Bentley, J., Odoulami, R. C., Pigot, A. L., & Trisos, C. H.: Risks to biodiversity from temperature overshoot pathways. Philosophical Transactions of the Royal Society B: Biological Sciences, 377(1857). https://doi.org/10.1098/rstb.2021.0394, 2022.

690 Mongwe, P., Long, M., Ito, T., Deutsch, C., and Santana-Falcón, Y.: Climatic controls on metabolic constraints in the ocean, Biogeosciences, 21, 3477–3490, <u>https://doi.org/10.5194/bg-21-3477-2024</u>, 2024.

Morée, A. L., Clarke, T. M., Cheung, W. W. L., & Frölicher, T. L.: Impact of deoxygenation and warming on global marine species in the 21st century. Biogeosciences, 20(12). <u>https://doi.org/10.5194/bg-20-2425-2023</u>, 2023.

Orr, J. A., Vinebrooke, R. D., Jackson, M. C., Kroeker, K. J., Kordas, R. L., Mantyka-Pringle, C., Van den Brink, P. J., de Laender, F., Stoks, R., Holmstrup, M., Matthaei, C. D., Monk, W. A., Penk, M. R., Leuzinger, S., Schäfer, R. B., Piggott, J. J.; Towards a unified study of multiple stressors: Divisions and common goals across research disciplines. Proceedings of the Royal Society B: Biological Sciences. <u>https://doi.org/10.1098/rspb.2020.0421</u>, 2020.

Oschlies, A.: A committed fourfold increase in ocean oxygen loss. Nature Communications, 12(1). https://doi.org/10.1038/s41467-021-22584-4, 2021.

Palomares, M. L., Stergiou, K. I., & Pauly, D.: Fishes in Databases and Ecosystems. Fisheries Centre Research Reports 14(4), 12(4), 2004.

Pfleiderer, P., Schleussner, C.-F., Sillmann, J: Limitied reversal of regional climate signals in overshoot scenarios. Environmental Research Letters, 3, 015005. <u>https://doi.org/10.1088/2752-5295/ad1c45</u>, 2023.

Pinsky, M. L., Selden, R. L., & Kitchel, Z. J.: Climate-Driven Shifts in Marine Species Ranges: Scaling from Organisms to
 Communities. Annual Review of Marine Science. <u>https://doi.org/10.1146/annurev-marine-010419-010916</u>, 2020.

Pörtner, H.O., (2010).: Oxygen-and capacity-limitation of thermal tolerance: a matrix for integrating climate-related stressor effects in marine ecosystems. Journal of Experimental Biology, 213(6), 881-893, <u>https://doi.org/10.1242/jeb.037523</u>, 2010.

Poloczanska, E. S., Burrows, M. T., Brown, C. J., Molinos, J. G., Halpern, B. S., Hoegh-Guldberg, O., Kappel, C. V., Moore, P. J., Richardson, A. J., Schoeman, D. S., Sydeman, W. J.: Responses of marine organisms to climate change across oceans. Frontiers in Marine Science. https://doi.org/10.3389/fmars.2016.00062, 2016.

Nature Editorial: Reversing climate overshoot. Nature Geoscience. https://doi.org/10.1038/s41561-023-01213-3, 2023.

710

715

Rogelj, J., Popp, A., Calvin, K. V., Luderer, G., Emmerling, J., Gernaat, D., Fujjmori, S., Strefler, J., Hasegawa, T., Marangoni, G., Krey, V., Kriegler, E., Riahi, K., van Vuuren, D. P., Doelman, J., Drouet, L., Edmonds, J., Fricko, O., Harmsen, M., Havlik, P., Humpenöder, F., Stehfest, E., Tavoni, M.: Scenarios towards limiting global mean temperature increase below 1.5 °C. Nature Climate Change, 8(4). https://doi.org/10.1038/s41558-018-0091-3, 2018.

Santana-Falcón, Y., Yamamoto, A., Lenton, A., Jones, C. D., Burger, F. A., John, J. G., Tjiputra, J., Schwinger, J., Kawamiya, M., Frölicher, T. L., Ziehn, T., Séférian, R.: Irreversible loss in marine ecosystem habitability after a temperature overshoot. Communications Earth & Environment, 4(1), 343. <u>https://doi.org/10.1038/s43247-023-01002-1</u>, 2023.

Schleussner, CF., Ganti, G., Lejeune, Q. Zhu, B., Pfleiderer, P., Prütz, R., Ciais, P., Fröilcher, T. L., Fuss, S., Gasser, T.,
Gidden, M. J., Kropf, C M., Lacroix, F., Lamboll, R., Martyr, R., Maussion, F., McCaughey, J. W., Meinshausen, M., Mengel,
M., Nicholls, Z., Quilcaille, Y., Sanderson, B., Seneviratne, S. I., Sillmann, Smith, C. J., Steiner, N. J., Theokritoff, E., Warren,

R., Price, J., Rogelj, J., Overconfidence in climate overshoot. Nature , 634, 366–373. <u>https://doi.org/10.1038/s41586-024-08020-9</u>, 2024.

Schmidtko, S., Stramma, L., & Visbeck, M.: Decline in global oceanic oxygen content during the past five decades. Nature, 542, 335. Retrieved from <a href="http://dx.doi.org/10.1038/nature21399">http://dx.doi.org/10.1038/nature21399</a>, 2017.

Schwinger, J., Asaadi, A., Steinert, N. J., & Lee, H.: Emit now, mitigate later? Earth system reversibility under overshoots of different magnitudes and durations. Earth System Dynamics, 13(4). <u>https://doi.org/10.5194/esd-13-1641-2022</u>, 2022.

Sumaila, U. R., Cheung, W., Dyck, A., Gueye, K., Huang, L., Lam, V., Pauly, D., Srinivasan, T., Swartz, W., Watson, R., Zeller, D.: Benefits of Rebuilding Global Marine Fisheries Outweigh Costs. PLOS ONE, 7(7), e40542. Retrieved from https://doi.org/10.1371/journal.pone.0040542, 2012.

Teh, L. C. L., & Sumaila, U. R.: Contribution of marine fisheries to worldwide employment. Fish and Fisheries, 14(1), 77–88. https://doi.org/10.1111/j.1467-2979.2011.00450.x, 2013.

Terhaar, J., Frölicher, T. L., Aschwanden, M. T., Friedlingstein, P., & Joos, F.: Adaptive emission reduction approach to reach any global warming target. Nature Climate Change, 12(12), 1136–1142. <u>https://doi.org/10.1038/s41558-022-01537-9</u>, 2022.

735 Zweng, M. M., Reagan, J. R., Seidov, D., Boyer, T. P., Antonov, J. I., Locarnini, R. A., Garcia, H. E., Mishonov, A. V., Baranova, O. K., Weathers, K. W., C. R. Paver, Smolyar, I. V.: World Ocean Atlas 2018 Volume 2: Salinity. NOAA Atlas NESDIS, 82(July), 2019.

740

730