RESEARCH ARTICLE

Ocean warming and novel species interactions boost growth and persistence of range-extending tropical fishes but challenge that of sympatric temperate species in temperate waters

Minami Sasak[i1](#page-0-0) | **Cristián J. Monaco[2](#page-0-1)** | **David J. Booth[3](#page-0-2)** | **Ivan Nagelkerke[n1](#page-0-0)**

1 Southern Seas Ecology Laboratories, School of Biological Sciences, the University of Adelaide, Adelaide, South Australia, Australia

2 IFREMER, IRD, Institut de Louis-Malardé, Univ Polynésie française, UMR-241 SECOPOL, Taravao, Tahiti, French Polynesia

³Fish Ecology Lab, Faculty of Science, University of Technology Sydney, Sydney, New South Wales, Australia

Correspondence

Ivan Nagelkerken, Southern Seas Ecology Laboratories, School of Biological Sciences, the University of Adelaide, Adelaide, SA, Australia. Email: ivan.nagelkerken@adelaide.edu.au

Funding information

Australian Research Council (ARC), Grant/Award Number: DP170101722 and DP230101932; NSW DPI, Grant/ Award Number: F94/696; University of Technology Sydney Animal Ethics Application, Grant/Award Number: 17-117; University of Adelaide Animal Ethics Application, Grant/Award Number: S-2018-107

Abstract

Aim: Climate change can have a broad range of impacts on the physiology and behaviour of animals. These effects can be mediated by the presence of other species in the community, but current forecasts of species responses to climate change largely ignore biological interactions. This is particularly true for novel interactions between range-extending and native species, as this is often considered as noise and excluded from predictive models. Here we simulate how a tropical range-extending and a local temperate fish species respond to the independent and combined effects of future ocean warming (RCPs 4.5 and 8.5) and novel ecological interactions in temperate ecosystems.

Location: East coast of Australia, along a ~2,000km latitudinal gradient in a global climate warming hotspot.

Taxon: *Abudefduf vaigiensis* (tropical) and *Atypicthys strigatus* (temperate) fishes.

Methods: We use a dynamic energy budget model to simulate the length growth (i.e., increases in body length of individuals over time) and population persistence of juveniles of a tropical and a temperate fish species that form mixed-species shoals, under different climate scenarios with and without the effects of novel ecological interactions.

Results: Our model forecasts that length growth of the juvenile tropical species will increase under ocean warming across subtropical to temperate regions. This increased length growth will be more drastic in temperate regions than in the subtropics, as winter warming will allow the tropical species to overwinter more frequently and show positive growth throughout the year. In contrast, warmer summer temperatures in the subtropics will likely exceed the optimal temperature of the juvenile temperate species at their trailing edge, resulting in reduced length growth under climate warming. Novel species interactions increased length growth of the juvenile tropical species but did not affect its winter or summer survival. In contrast, novel species interactions with

This is an open access article under the terms of the [Creative Commons Attribution](http://creativecommons.org/licenses/by/4.0/) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2024 The Author(s). *Journal of Biogeography* published by John Wiley & Sons Ltd.

munities (Alexander et al., [2015](#page-11-1)). tropical species were forecast to reduce length growth of the juvenile temperate **KEYWORDS** climate change, dynamic energy budget, novel species interactions, ocean warming, range shifts, vagrant species

1 | **INTRODUCTION**

Climate change is reshuffling biological communities, as species track changing environmental temperatures at different rates (Chen et al., [2011](#page-11-0); Poloczanska et al., [2013](#page-13-0)). Notably, warming facilitates the extension of biogeographic ranges of warm-adapted species towards ecosystems and higher latitudes that were previously unsuitable for them. This has resulted in shifts in biodiversity along temperature gradients. For instance, warming-induced range shifts have caused declines in species diversity at lower latitude while leading to increases at higher latitudes, where tropical range-extending species mix with temperate species (Worm & Lotze, [2021](#page-14-0)).

In the temperate recipient ecosystems where tropical rangeshifting species interact with resident temperate species, different fish species will likely respond to ocean temperature changes differently due to each species having its own optimal temperature range. Because fish species originating from warm regions have higher optimal temperatures compared to those in cooler regions (Payne et al., [2016\)](#page-13-1), ocean warming at higher latitudes will likely favour those that are shifting from the tropics to cooler temperate waters compared to those that are resident to higher latitudes. Given that changes in ambient temperature have a strong influence on the metabolic rate of fish species (Sinclair et al., [2016](#page-14-1); Volkoff & Rønnestad, [2020](#page-14-2)), changes in energy demand associated with temperature increases can lead to shifts in energy demand and growth rate of fish species (Volkoff & Rønnestad, [2020](#page-14-2)). Hence, future ocean warming at higher latitudes will likely enhance growth rate of the range-shifting species but decline that of the residents. In addition to direct impact of climate change on the physiological performance, arrival of range-shifting species can challenge resident species via competition (Zeidberg & Robison, [2007\)](#page-14-3) and predation (Hollebone & Hay, [2008](#page-12-0); Vergés et al., [2014](#page-14-4)), or by modifying local habitat composition (Ling, [2008](#page-14-5); Whitney & Gabler, 2008), thus threatening to alter the structure of the communities they occupy (Kordas et al., [2011](#page-12-1); Williams & Jackson, [2007](#page-14-6)). As such, a better understanding of the combined effects of climate change-related stressors and novel species interactions is key to forecasting the

species.

Main Conclusions: Our study suggests that for some coastal fish species future warming will likely reverse body size dominance between temperate and tropical fish species, with increased novel interactions in temperate ecosystems (due to range extensions) but decreased novel interactions in the subtropics (due to range contractions). Novel species interactions and warming effects on body size and species survival are likely to reshuffle temperate fish communities and their competitive interactions.

ecological consequences of poleward range shifts on recipient com-

The ecological success of species depends both on biotic and abiotic elements of the environment. However, current forecasts of species responses to climate change still largely exclude species interactions (Dolci & Peruzzi, [2022](#page-12-2); Gillard et al., [2017](#page-12-3); Guisan et al., [2006](#page-12-4)), and especially competition (Araújo & Luoto, [2007](#page-11-2)). Some models exist that include predator–prey interactions (e.g., aquatic taxa, Monaco et al., [2016;](#page-13-3) Moullec et al., [2019](#page-13-4), terrestrial taxa, Aryal et al., [2016](#page-11-3); Bastille-Rousseau et al., [2018](#page-11-4)). Although some studies model species' future physiological performance and include competitive interactions, these studies are uncommon and largely limited to species with limited ability to move (e.g., mussels: Monaco & McQuaid, [2019](#page-13-5); trees: Caplat et al., [2008;](#page-11-5) Liang et al., [2017](#page-13-6)). This leaves a critical knowledge gap concerning aquatic mobile taxa as range extensions by mobile fish species are increasingly documented (Osland et al., [2021](#page-13-7); Pinsky et al., [2020;](#page-13-8) Poloczanska et al., [2013](#page-13-0); Vergés et al., [2014](#page-14-4)). These species have the potential to compete with resident species due to their shared utilization of available resources, which may significantly impact resident species given their susceptibility to environmental change (Manes et al., [2021](#page-13-9)). Furthermore, warming-related range shifts (Gervais et al., [2021](#page-12-5); Hyndes et al., [2016](#page-12-6); Johnson et al., [2011](#page-12-7)) suggest that increasing ocean temperatures will likely accelerate the rate of range expansion. As such, predicting how novel species interactions and future climate will affect the persistence and physiological performance of species is an urgent outstanding question.

A common approach to predicting species responses to a changing climate is the use of correlative niche models, such as species distribution models. These models are mathematical representations of the environmental niche of a species, built based on the conditions (e.g., temperature, salinity, chlorophyll *a*) they experience across their distribution range. These models can provide estimates of the future distribution of the species by modifying the habitat temperatures based on climate change projections. To capture the thermal niche of a species (Peterson & Vieglais, [2001](#page-13-10)), these correlative models often assume that the distribution range corresponds to its entire

native range (Araújo et al., [2005](#page-11-6); Araújo & Pearson, [2005](#page-11-7); Guisan & Zimmermann, [2000](#page-12-8)). However, it is not uncommon to remove individuals that occur outside of the species native range ("vagrants") from correlative models because they are considered as a source of error (e.g., Cooper & Soberón, [2017](#page-12-9); D'Amen & Azzurro, [2019](#page-12-10); Rose et al., [2016\)](#page-13-11). Filtering these individuals consequently removes information regarding the novel interactions occurring between a vagrant and local species as well as their novel climate niche. Although comparing the analyses with a filtered niche across a range of percentiles (e.g., from 75% to 100%) allows researchers to understand the impact of removing novel environments (Guisan et al., [2014](#page-12-11)), when model calibrations are performed on such data, predictions beyond present-day conditions can consequently be compromised due to calibration ignoring novel conditions (Lee-Yaw et al., [2022;](#page-12-12) Liu et al., [2020](#page-13-12); Zurell et al., [2012](#page-14-7)). These drawbacks in niche modelling can be accommodated by using either shape-constrained GAMs (Citores et al., [2020](#page-12-13)) or mechanistic models designed to explicitly account for direct physiological and indirect behavioural responses to climate change-related drivers. For example, the Dynamic Energy Budget (DEB) model framework (Kooijman, [2010](#page-12-14)) uses an organisms' physiology to identify suitable environmental conditions of the species, instead of reconstructing species niches from their geographical distribution (Kearney et al., [2008](#page-12-15)). As this mechanistic model uses the species' fundamental niche (as opposed to the realized niche in species distribution models), more robust predictions beyond their native ranges are possible.

DEB theory is a powerful framework to describe an organisms' energy metabolism throughout their life cycle (Marques et al., [2018](#page-13-13); Van der Meer, [2006\)](#page-14-8). Variations of the DEB model has been used to model the physiological performance of species across geographical scales, including predictions under future climate change scenarios (bivalves: Thomas & Bacher, [2018](#page-14-9); Monaco & McQuaid, [2019](#page-13-5)). DEB models have been developed to accommodate competitive interactions based on changes in the individual's behaviour (Beale & Lennon, [2012](#page-11-8)); however, these model variants have not been explored in a context of species range shifts due to climate change.

Here we modelled the physiological performance, based on survival and growth, of a common range-extending coral-reef fish, the Indo-Pacific sergeant major *Abudefduf vaigiensis* (hereafter "tropical species"), and a shoaling local temperate fish, Australian mado *Atypichthys strigatus* (hereafter "temperate species"), in a warming hotspot in temperate SE Australia. *A. vaigiensis* occurs from tropical to warm-temperate waters (from 36°N to 39°S, Froese & Pauly, [2000](#page-12-16)) and their breeding population has been observed at a latitude as high as 28.7°S based on western Australian records (Pearce et al., [2016](#page-13-14)). Note that observations of breeding behaviour at such high latitudes in eastern Australia are not available. This tropical species feeds on zooplankton, benthic algae, and small invertebrates (Allen & Erdmann, [2012](#page-11-9); Sasaki et al., [2024](#page-14-10)). *A. strigatus* is endemic to temperate waters of eastern Australia (Bray, [2023](#page-11-10)). Similar to the tropical species, *A. strigatus* feeds on zooplankton and benthic organisms (e.g., algae and invertebrates) as well as parasitic organisms

(e.g., caligoids and monstrilloids) (Glasby & Kingsford, [1994](#page-12-17); Sasaki et al., [2024](#page-14-10)). The tropical *A. vaigiensis* is one of the most abundant vagrant species colonizing climate-warming hotspots (Ridgway, [2007\)](#page-13-15) of temperate Australia during late summer (Booth et al., [2018;](#page-11-11) Monaco et al., [2020](#page-13-16)). Juveniles of this tropical species have been observed to school with juveniles of *A. strigatus* (Smith et al., [2017\)](#page-14-11) along with Eastern hula fish (*Trachinops taeniaus*) (Smith et al., [2017\)](#page-14-11) and stripey (*Microcanthus strigatus*, Coni et al., [2021](#page-12-18)). Previous studies showed mixed types of interaction between *A. vaigiensis* and *A. strigatus*. For instance, the interaction with juveniles of the temperate species in temperate Australia (i.e., novel environment for the tropical species) led to enhanced growth in *A. vaigiensis* towards winter (Smith et al., [2017](#page-14-11)), showing a positive interaction outcome. By contrast, *A. strigatus* interacted aggressively and prevented *A. vaigiensis* from accessing food during a feeding experiment in the field (Coni et al., [2021](#page-12-18)), showing a negative interaction. Despite the mixed results, overlaps in diet between *A. vaigiensis* and *A. strigatus* (e.g., calanoids and cyclopoids, Glasby & Kingsford, [1994](#page-12-17); Sasaki et al., [2024](#page-14-10)) make the juveniles of tropical *A. vaigiensis* potential competitors to the juveniles of temperate species. Furthermore, increases in the abundance of range-shifting tropical species may not be favourable to the local species due to a positive relationship between the abundance of range-shifting tropical species and their feeding rate (Coni et al., [2021](#page-12-18)). As such, we used these two species as model organisms to study the effects of novel species interactions due to climate change.

Currently, vagrant populations of this tropical species in temperate Australia (leading edge of the species distribution) still suffer from very high (10% overwinter ratio based on 18-year survey, Booth et al., [2018](#page-11-11)) winter mortality because temperatures drop below their thermal minimum. As such, novel interactions between the tropical and the temperate species are restricted to the warmer parts of the year and largely cease in winter before the following summer recruitment event occurs. However, with climate changedriven temperature increases, their survival through winter is expected to increase (Figueira et al., [2009](#page-12-19); Figueira & Booth, [2010;](#page-12-20) Monaco et al., [2020](#page-13-16)), extending the duration of these novel interactions.

In contrast to tropical range extenders, cool-adapted temperate species, particularly the populations located at their warm range trailing edge, are expected to experience reduced physiological and behavioural performance with increasing and potentially stressful levels of warming (Donelson et al., [2019](#page-12-21)). As such, cool-adapted species are expected to suffer from either reduced growth or increased mortality, or shift to more suitable habitats as seen in several taxa (terrestrial organisms: Parmesan, [2006](#page-13-17); marine organisms: Pinsky et al., [2020](#page-13-8)). If their populations decline at their trailing edges, this might also reduce the magnitude of novel interactions and possible minimize negative effects on tropical range extenders. Thus, future warming is expected to affect both the physiological condition of temperate fish species and the magnitude of their potential ecological interactions with range extending tropical species.

13652699, 2024, 11, Downloaded from https://onlinelibrary.wiley.com/doi/10.1111/jbi.14983 by Ifremer Centre Bretagne Blp, Wiley Online Library on [21/10/2024]. See the Terms and Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons License

and Conditions

sdnup:

library.wiley

conditions š

Wiley Online Library

for rules

use; OA

os are

erned by the

applicable Creative

Common

Online Library on [21/10/2024]. See the Terms

3652699, 2024, 11, Downloaded from https:

kша

wiley.com/doi/10.1111/jbi.14983 by Ifremer Centre

Bretagne Blp,

, Wiley

Here, we modelled the thermal physiological performance of tropical and temperate juvenile fishes in Australia to understand how future warming and novel species interactions might impact communities. We hypothesised that future ocean warming will enhance the growth rate and survival duration of juveniles of tropical species compared to temperate species in temperate regions. Additionally, we hypothesised that the presence of novel interactions has a mediating effect (either positively or negatively) on the physiological responses (i.e., growth rate and survival duration) of our model species. For this, we parameterised the DEB model for a sympatric juvenile tropical and temperate fish species, using life-history traits (e.g., body size, age, and reproductive output (obtained from published data, Table [S1\)](#page-15-0)) and species thermal sensitivities. Importantly, we included novel ecological interactions based on food ingestion rates and growth rates in the presence and absence of novel interactions based on controlled aquarium experiments. Our model allowed us to forecast the temporal persistence and growth of the two species under present-day and two future ocean warming scenarios (i.e., RCP 4.5 and RCP 8.5) with and without novel species interactions. Because the DEB model is not spatially explicit, the current simulation was limited to latitudes between 27.5 to 43.3°S along the southeast coast of Australia where two species are known to occur and where novel ecological interactions due to climate warming are currently most common.

2 | **MATERIALS AND METHODS**

2.1 | **Future climate projections**

We used a dynamic energy budget (DEB) model to parameterise parameters to simulate the physiological change of juvenile tropical and temperate species in Australia under future warming scenarios with and without novel interactions. The DEB model description including diagrams showing the model differences between non-stressful and stressful conditions (Figure [S1\)](#page-15-0), parameterisation, use of individual-based DEB model, and testing for model fit is found in the [Appendix](#page-15-1).

The simulations were limited to the juvenile stages as we were interested in growth of the tropical species when they recruit in temperate regions during the summer. This means that we excluded the larval period and considered fish performance from settlement stage sizes, i.e., >1.21 and 1.20 cm for the tropical and the temperate species, respectively. We restricted our geographical range for simulation of growth to latitudes 27.5–43.3°S along the southeast coast of Australia, covering the known occurrence range of the vagrant population of the tropical species (Atlas of Living Australia, [2021a](#page-11-12)) and the known distribution of the temperate species (Atlas of Living Australia, [2021b](#page-11-13)). The latitudinal ranges used in the simulation cover several climate zones including subtropics (from 27.5 to 30.5°S), warm-temperate (from 31.5 to 34.5°S), mild-temperate (from 35.5 to 38.5°S) and cool temperate (from 39.5 to 43.3°S) areas. The sites

used for simulation were chosen at every one degree of latitude from the lowest latitude of 27.5°S to the highest at 43.3°S, resulting in a total of 11 and 16 sites for the tropical and the temperate species, respectively.

Within these settings, we simulated growth in body size and development in maturity of the two species for 1 year, from the first of January to the end of December using the parameters estimated (Table [S2](#page-15-0)) in the DEB_IBM-model version 3.1 (Martin et al., [2012](#page-13-18)) in NetLogo 6.1 (Wilensky, [1999](#page-14-12)). Note that we did not account for the density-body size relationship (Cyr, [2000](#page-12-22); Cyr et al., [1997\)](#page-12-23). We selected January because both species have been observed to start recruiting during this month in SE Australia (Table [S3](#page-15-0)). Note that January recruitment of two species at the same site does not always occur (Table [S3\)](#page-15-0), but we still observed the two species co-occurring at the same site in temperate Australia (Figure [S2\)](#page-15-0). At the end of each simulation, we obtained the total number of months in which individuals of each species had sufficient energy to grow ("persistence duration") and the total standard length ("length growth"). Note that persistence duration reflects a temporal duration from the beginning of a simulation (i.e., January) until a month when the level of energy reserves falls below the scaled length (Martin et al., [2010](#page-13-19)). For the total standard length, we sum length growth of an individual across persistence duration.

Two variables that we used in the simulation were monthly averaged sea-surface temperature (SST) (current and future year 2100) and presence/absence of species interaction. For the SST data, we retrieved satellite-derived SST for the year 2018 from Australia's Integrated Marine Observing System (IMOS) (IMOS, [2019](#page-12-24)) and projected SST for the year 2100 from WDC CLIMATE ([2019](#page-14-13)). The latter was derived using the Australian Community Climate and Earth System Simulator (ACCESS) under the Coupled Model Intercomparison Project Phase 5 (CIMP5). We used two Representative Concentration Pathways (RCP), the intermediate RCP 4.5 and the high RCP 8.5, representing greenhouse gas emissions leading to radiative forcing in 2100 stabilizing at 4.5 and 8.5 W m−2, respectively. On average, the ocean temperatures across all simulated sites increases by 1.10 and 3.42°C in 2100, relative to the annual average ocean temperature in 2018, under RCP 4.5 and RCP 8.5, respectively (Figure [S3](#page-15-0)). Of all simulated latitudes, latitude 28.5°S showed the highest yearly average ocean temperature in all scenarios (Figure [S3\)](#page-15-0) and scored the highest summer temperature of 30.63°C in 2100 under RPC 8.5.

We simulated the effect of presence/absence of novel species interactions using the feeding rates of individuals as estimated in the aquarium experiment: *f_paired* (Table [S2](#page-15-0)) and *f_single* (Table [S2\)](#page-15-0) were used for the presence and absence of the interaction, respectively. Novel interactions occurred only when the months of persistence overlapped between two species ("duration of the novel interaction"). We did not simulate the effect of species interactions at latitudes ≥37.5°S in the current study as it is uncertain if the tropical species will cross the Bass Strait to reach the island of Tasmania by 2100.

Comparison of persistence duration and length growth between current and future scenarios was used to understand the effect of future warming and species interactions on fish physiological performance.

3 | **RESULTS**

3.1 | **Model fit**

We found that the model with interactions predicted the maximum standard length of both species slightly better than the model without interactions, with a mean absolute percentage error of −0.036 (Figure [S4b](#page-15-0)) compared to 0.067 (Figure [S4a\)](#page-15-0), respectively. These error rates were slightly worse than the average relative error estimated across 500 DEB models (the median mean relative error of <0.1, Marques et al., [2018](#page-13-13)). The correlations between predicted and observed maximum standard length improved from 0.524 (Figure [S4c](#page-15-0)) to 0.724 (Figure [S4e](#page-15-0)) for the tropical species and from 0.069 (Figure [S4d\)](#page-15-0) to 0.094 (Figure [S4f\)](#page-15-0) for the temperate species as species interaction was included in the model. The mean absolute percentage error of model fit further improved to 0.024 (Figure [1a](#page-4-0)) as we selected the model (with or without the interaction) based on the lowest absolute percentage error between the observed and predicted size. The correlations between predicted and observed maximum standard length improved to 0.901 for the tropical species (Figure [1b](#page-4-0)) and 0.234 for the temperate species (Figure [1c](#page-4-0)).

3.2 | **The effect of climate change on body length growth (without novel species interactions)**

Increasing ocean warming is predicted to boost length growth of the juvenile tropical species by an average 41.5% under RCP 4.5 and 58.4% under RCP 8.5, respectively (Figure [2a](#page-5-0)). The rate of length growth increase in the juvenile tropical species is more extensive at warm- to mild-temperate (32.5–37.5°S) compared to subtropical to warm-temperate (27.5–31.5°S) latitudes: 70.5% vs 6.7% (RCP 4.5) and 97.8% vs 10.9% (RCP 8.5), respectively (Figure [2a](#page-5-0)). Nevertheless, largest absolute lengths were still accomplished at subtropical to warm-temperate latitudes (27.5–31.5°S) (Figure [2c](#page-5-0)).

In contrast, future warming is projected to be less favourable for the juvenile temperate species. Under RCP 4.5, the rate of change in juvenile length growth was on average +3.2, +15.0 and −21.8% at the trailing (i.e., subtropics between 27.5 and 30.5°S), core (i.e., warm- to mild-temperate between 31.5 and 37.5°S) and leading edges (i.e., mild- to cool-temperate between 38.5 and 43.3°S), respectively (Figure [2b](#page-5-0)). Under RCP 8.5, juvenile length growth is predicted to drop by 36.8% at the trailing edge, but to increase by 14.3% at the core and by 74.4% at the leading edges, respectively (Figure [2b](#page-5-0)). Under both climate change scenarios, largest fish sizes are forecast to shift in occurrence from the current trailing edge to the core range (Figure [2d](#page-5-0)).

Under present-day ocean temperatures, average body size of the juvenile tropical species is smaller than that of juvenile temperate species by 8.0%–42.5% at subtropical to mild-temperate latitudes (i.e., 27.5–37.5°S). However, in year 2100 under RCP 8.5, the juvenile

FIGURE 1 Model performance based on absolute percentage error between observed maximum length and predicted length for model with lowest absolute percentage error (i.e., "Best models") (a). Mean absolute percentage error (MAPE) was calculated and shown at top left corner of each plot. Correlations between simulated standard length and observed maximum length for the tropical (b) and temperate species (c). Slope values (*b*) and their significance (*) from a 1:1 relationship (hatched lines) are provided. Shaded areas show 95% confident intervals.

FIGURE 2 Simulated growth length using a model without interaction under a current ("Current") and two future ("RCP4.5" and "RCP8.5") climate change scenarios for the tropical species (a, c) and the temperate species (b, d). The results were expressed as percentage difference ("% difference standard length") in final standard length (SL) between a current and future scenario (calculated as 100 \times (future SL – current SL)/current SL) (a, b), and as absolute standard length ("absolute standard length") (c, d). Latitudinal position is shown on the *y*-axis. The current simulation was performed at sites where the species are known to occur along the east coast of Australia (grey area=Australian continent).

tropical species is predicted to reach body lengths larger than that of the juvenile temperate species by 5.8%–15.8% at subtropical to warm-temperate latitudes (27.5–31.5°S) (Figure [2c,d\)](#page-5-0) but not at colder latitudes (32.5–37.5°S). On average across all latitudes simulated, the length difference between the juvenile tropical and temperate species will be reduced from 40.0% under current conditions to 16.0% in the year 2100 under RCP 8.5, and to 21.0% under RCP 4.5.

3.3 | **The effect of climate change on population persistence (without novel species interactions)**

Our model forecasts that the local population persistence of the juvenile tropical species will increase at its cold range edge under RCP 4.5 and 8.5, compared to present-day conditions. Under present-day temperatures, full population persistence (i.e., 365 days) of the juvenile tropical species is only observed in subtropical waters (i.e., 27.5–30.5°S), whereas under RCP 4.5 and RCP 8.5 the juvenile tropical species is predicted to persist throughout the year in both warm- and mild-temperate latitudes (i.e., up to 37.5°S; Figure [3a](#page-6-0)). In contrast, the model for the juvenile temperate species predicts little to no changes in population persistence across its leading, core and trailing edges (27.5–43.3°S) under either RCP 4.5 or RCP 8.5, except for a reduction from 365 to 50 days at a single subtropical latitude (28.5°S) under RCP 8.5 (Figure [3b](#page-6-0)) where a highest summer temperature of 30.63°C is forecast.

Idiosyncratic changes in population persistence in response to ocean warming will alter the time window for the two species to interact. At the subtropical latitude of 28.5°S under RCP 8.5, the time window for novel interactions will drop from 365 to 50 days. However, at warm- to mild-temperate latitudes between 32.5 and 37.5°S, the time window for novel interactions between the two species will increase from an average of 189.5 days under current conditions (range between 130–240 days) to 365 days under both RCP 4.5 and RCP 8.5 scenarios (Figure [3c](#page-6-0)).

3.4 | **The effect of novel ecological interactions on body length growth and population persistence**

Including novel interactions in the model did not alter the population persistence of either species under either climate change scenario at any latitude studied (Figure [S5a,b](#page-15-0)). In contrast, inclusion of species interactions altered the simulated body length increase of both species. Length of the juvenile tropical species increased when interacting with the juvenile temperate species by an average +25.8, +25.1 and +24.9% under present-day, RCP 4.5 and RCP 8.5 scenarios, respectively (Figure [4a](#page-7-0)). In contrast, that of the juvenile temperate species was reduced under novel interactions by an average −5.0, −5.9 and −5.3% under present-day, RCP 4.5 and RCP 8.5 scenarios, respectively (Figure [4b](#page-7-0)).

FIGURE 3 Population persistence for the tropical species (a) and the tempeate species (b) under the current ("2018 Current") and two future ("2100 RCP4.5" and "2100 RCP8.5") climate change scenarios based on a model without species interactions. Latitudinal position is shown on the y-axis and population persistence in days on the *x*-axis. (c) Duration of novel interactions indicates the total months for which the population persistence of both species overlapped. Colours used for different scenarios: Current (blue), RCP 4.5 (green) and RCP 8.5 (pink) scenarios, respectively.

The model further predicted that the effect of novel species interactions on the change in length growth in both species was greater at subtropical to warm-temperate latitudes (27.5–31.5°S) than warm- to mild-temperate latitudes (32.5–37.5°S) under present-day conditions (Figure [4a,b](#page-7-0)). This latitudinal pattern in novel interactions effect was observed in the body size of juvenile tropical species. Inclusion of novel interactions in the model simulated the body size of juvenile topical to be larger than that of juvenile temperate species by 7.8% to 26.3% at subtropical to warm-temperate latitudes (27.5–31.5°S) but smaller by 24.9% to 39.9% at warm-temperate to mild-temperate latitudes (32.5–37.5°S) (Figure [4c,d](#page-7-0)). Under RCP 4.5 scenario, this latitudinal decline effect is predicted to remain for the

 -2.5

 -5.0

 -7.5

FIGURE 4 Changes in growth length between a model with interaction ("with") and without interaction ("no") under a current ("Current") and two future ("RCP4.5" and "RCP8.5") climate change scenarios for the tropical species (a) and the temperate species (b). The results were expressed as percentage difference ("% difference standard length") in final standard length (SL) between a model with interaction and without interaction (calculated as 100 × (SL without interaction – SL with interaction)/SL with interaction) (a, b). The direction of body length change is indicated by different colours: Warm red colours = positive effects, cool blue colours = negative effects, and the magnitude of change in length is indicated by the colour gradients. Simulated absolute standard length using a model with interaction for the tropical species (c) and the temperate species (d). Latitudinal position is shown on the *y*-axis. The current simulation was performed at sites where the species are known to occur along the east coast of Australia (grey area=Australian continent).

juvenile tropical species, but increase at mild-temperate latitudes (27.5–37.5°S) for the juvenile temperate species. Under RCP 8.5, the change in length growth due to species interactions will become

similar across latitudes for both species, except at latitude of 28.5°S for the juvenile temperate species (Figure [4b](#page-7-0)). The juvenile tropical species is predicted to reach body lengths larger than that of the juvenile temperate species by 3.8% to 5.6% at warm-temperate latitudes (32.5–34.5°S) (Figure [4c,d](#page-7-0)) under RCP 4.5 and by 10.7% to 12.3% at mild-temperate latitudes (35.5–37.5°S). The above patterns closely matched those for the duration for which the two species cooccur (i.e., novel interactions; Figure [3c](#page-6-0)).

4 | **DISCUSSION**

To predict the impacts of climate change on juvenile fish growth and their population persistence, we parameterised and used the Dynamic Energy Budget model addressing two factors: speciesspecific thermally-dependent physiological rates (mediated by abiotic factors) and the effect of novel species interactions (mediated by biotic factors). When simulating the impact of future warming without novel species interactions, our model showed that the juvenile tropical species, *Abudefduf vaigiensis* will likely benefit from future warming in terms of population persistence and magnitude of body length growth at its leading range edge in temperate ecosystems. In contrast, the juvenile temperate species, *Atypicthys strigatus*, is predicted to experience reduced growth at their trailing edge, and even suffer from inability to sustain positive growth during future summer temperatures (based on monthly average temperatures) under an RCP 8.5 scenario. When including the effects of novel species interactions, our model forecasts that these interactions will likely influence the length growth positively for the tropical species but negatively for the temperate species.

4.1 | **Species responses to ocean warming**

Temperature had significant effects on the population persistence and body length growth in both species within the simulated geographical range. However, due to differences in thermal sensitivity between the tropical and temperature species, future warming will act on them differently. More specifically, warmer winters will improve the yearly growth performance of the juvenile tropical species at its leading cool range edge, whereas warmer summers will reduce that of the juvenile temperate species at its trailing warm range edge.

Increased population persistence and somatic growth in the juvenile tropical species across simulated latitudinal range in the year 2100 agrees with the current understanding that: (1) growth potential of *Abudefduf vaigiensis* is not being fulfilled under current cool temperatures, but that: (2) under warming waters, physiological performance including overwintering (Figueira & Booth, [2010](#page-12-20)) and growth (Kingsbury et al., [2020](#page-12-25)) will improve. Under present-day conditions, the abundance of juvenile *A. vaigiensis* drops with decreasing ocean temperatures at increasing latitudes (Figueira & Booth, [2010](#page-12-20)) and only a fraction of the fish survive through winter in warm- and mildtemperate waters (Booth et al., [2018](#page-11-11)). Considering the predicted future winter warming, our results suggest that the potential for *A. vaigiensis* colonization along the temperate southeast coast of Australia could greatly increase if greenhouse gas emissions follow the trajectory predicted by RCP 4.5 or RCP 8.5. Although we did not simulate

reproduction in the current study, the ability to overwinter and attain maximum (reproductive) body size suggests the potential for *A. vaigiensis* populations to increase with subsequent permanent establishment in most temperate SE Australian regions by the year 2100.

We found that summer ocean temperatures forecast for the year 2100 under an RCP 8.5 scenario will likely exceed the optimal temperature for the temperate species, *Atypichthys strigatus* at its trailing edge of distribution (i.e., at 28.5°S). This appears to result in reduced growth or even in losing thermally suitable habitat at subtropical latitudes. Our prediction is consistent with previous findings that populations of temperate species located at their warm range edge experience a reduction in growth (Neuheimer et al., [2011](#page-13-20)) or extirpation (Smale & Wernberg, [2013](#page-14-14); Wernberg et al., [2013](#page-14-15)). As growth is aerobic, limited oxygen supply to match required oxygen demand (i.e., aerobic scope) due to thermal stress likely constraints growth (Pörtner & Knust, [2007\)](#page-13-21). Considering that monthly averaged ocean temperature under RCP scenario 4.5 and 8.5 was sufficient to compromise growth performance of the juvenile temperate species at its trailing edges, further impairment of growth can be expected under extreme environmental conditions such as marine heat waves (Van der Walt et al., [2021](#page-14-16)). Losing thermally suitable habitat at their warm range edge is of great concern for this species as there is no landmass between Tasmania and Antarctica. This may lead to the temperate species having to contract its distributional range to a narrow band of latitudes, as also predicted for other species will eventually run out of suitable climate space (Freeman et al., [2018;](#page-12-26) Whitfield et al., [2016\)](#page-14-17). However, the temperate species may be able to extend its ranges to deeper colder waters in response to climate stress as reportedly previously for other marine species (Nye et al., [2009;](#page-13-22) Perry et al., [2005](#page-13-23); Pinsky et al., [2013](#page-13-24)), with *Atypichthys strigatus* shown to occur to depth of 30 m (Bray & Gomon, [2011](#page-11-14)). Nonetheless, the effect of future warming on the growth rate for this species will still likely be adverse at its trailing edge.

In the current study, we utilized average sea surface temperatures in our simulations; therefore, our results may differ under extreme weather conditions. Global climate change is projected to increase the frequency of extreme weather events (Seneviratne et al., [2021](#page-14-18)), and some studies have reported that such events can lead to population declines and local extirpations (Maxwell et al., [2019](#page-13-25)). While long-term exposure to warmer temperatures has been shown to increase ectotherms' critical thermal maximum (upper thermal tolerance of species) (Stewart et al., [2023](#page-14-19)) and improve aerobic performance at higher temperatures (Donelson et al., [2011](#page-12-27)), other research indicates that the capacity for maximum thermal tolerance remains relatively conserved even after acclimation to warmer water over multiple generations (Sandblom et al., [2016\)](#page-14-20). If our temperate species are unable to enhance their maximum thermal tolerance in the future, their persistence duration is likely to be reduced during extreme weather events.

4.2 | **Species responses to novel ecological interactions**

The model predicted that the presence of novel interactions affected the growth rate positively in the juvenile tropical species but

negatively in the juvenile temperate species because novel interaction increased functional response of the tropical species but decreased that of the temperate species.

The current modelled responses of novel interactions on juvenile body growth might differ from those in the wild because factors such as prey diversity, prey availability, mortality due to predation, and fish abundances were not accounted for in the laboratory experiment. For example, prey diversity is shown to positively correlate with the degree of food resource partitioning between species (Holbrook & Schmitt, [1989](#page-12-28); Sánchez-Hernández et al., [2017](#page-13-26)). Indeed, in the field where multiple prey species are available, our studied species did not feed on the exact same prey species (Kingsbury et al., [2019](#page-12-29)). Unlike in nature, the two species were fed one prey species (*Artemia*) in the aquarium experiment, which could have forced two species to interact more often and induced a stronger magnitude of novel interactions (i.e., due to altered feeding conditions). Furthermore, currently available prey species might not be present in the future because of a mismatch in the rate of range shift between predatory fish and zooplankton under warming conditions. For example, copepod species exhibited poleward range shift with a rate of 99 km per decade (Chivers et al., [2017](#page-11-15)), compared to a rate of poleward extension of 277.5 km per decade for bony fish in general (Poloczanska et al., [2013](#page-13-0)). The abundance of species has also been reported to affect feeding interactions of our species. A field study revealed that numerically abundant temperate species interfered the tropical species from accessing prey by forming a barrier between the supplied food source and the tropical species (Coni et al., [2021](#page-12-18)). Likewise, Monaco et al. ([2020](#page-13-16)) demonstrated that density feedback played important role in the growth of *A. vaigiensis* on earlier life stage (i.e., settler and juvenile). Although abundance of immigrants is usually lower than that of the resident fishes (de Souza & Dos Santos, [2023](#page-12-30)), these previous studies show the importance of fish abundance on the performance rate of the tropical species. As we used one individual of each species in our experiment, abundancerelated modification of feeding behaviours was not included in our simulation. Although obtaining species feeding responses using a laboratory experiment was crucial for estimating parameters that are affected by the presence/absence of novel interactions, the ability of the aquarium experiment to replicate fish response in the nature is limited. Consequently, the results from our model should be interpreted according to our aquarium setting (limited resources and equal fish abundances).

4.3 | **Overall implications**

Our results suggest that some coastal fish species will face drastic changes in their juvenile length growth and population persistence under ocean warming expected to occur at the end of the century in temperate ecosystems if greenhouse gas emissions continue to increase. In a temperature mixing zone where tropical and temperate species co-occur, ocean warming will have opposing

effects on species depending on their thermal affinities. This could subsequently alter present-day species interactions with ensuing effects on fish community compositions.

The contrasting growth responses shown by the two species in the subtropics to warm-temperate regions under RCP 8.5 will likely reverse the current body size advantage of the temperate species in the future. Our model forecasts that the juvenile tropical species could grow larger than the juvenile temperate species if the two species recruit at the same time at these latitudes under RCP 8.5 (but not RCP 4.5). The projected reversal in body size or reduction of inter-species body size differences might reshuffle the competitive interactions between tropical and co-schooling temperate fishes as body size has been shown to strongly mediate species interactions (e.g., Wissinger, [1988](#page-14-21); Woodward & Hildrew, [2002](#page-14-22)). For example, in South America the number of attacks made by competitively superior invasive Nile tilapia (*Oreochromis niloticus*) towards native pearl cichlids (*Geophagus brasiliensis*) was greater when the size of tilapia was larger. However, tilapia attacked cichlids significantly less frequently when the size of cichlids was 30% or 50% larger than that of tilapia (Sanches et al., [2012](#page-13-27)). In our aquarium experiment, the temperate species was on average 21% larger than the tropical species across 16 of 20 mixed-species pairs. Thus, the magnitude of novel species interactions in the experiment was largely based on the case where the temperate species was larger than the tropical species. Alternatively, differences in body size between the tropical and temperate species might facilitate segregation in species interactions because the strength of species interactions can sometimes be higher between similar-sized individuals than different-sized individuals (Woodward & Hildrew, [2002](#page-14-22); Kohda et al., [2008](#page-12-31)).

Warming winter will transform warm- to mild-temperate ecosystems to become more suitable for the juvenile tropical species, likely increasing the temporal duration of novel interactions between the temperate and the tropical species at higher latitudes under RCP 4.5 and 8.5. Relaxing winter environmental filtering with ongoing warming will likely facilitate other vagrant species that are known to occur along the east coast of Australia to overwinter as well and create novel interactions with a range of temperate fishes. For example, vagrant damselfish species from the genus *Abudefduf*, *Pomacentrus* and *Stegastes* displayed minimum temperature thresholds for winter survival of approximately 17°C, similar to our species *A. vaigiensis* (Figueira & Booth, [2010](#page-12-20)). The same applies to vagrant butterflyfish species from the genus *Chaetodon*, which displayed minimum temperature thresholds of a few degrees higher than *A. vaigiensis* (Figueira & Booth, [2010](#page-12-20)). Despite such predictions, we still have limited understanding as to how prolonged interactions of surviving tropical species during temperate winters will affect novel species interactions in more realistic communities (Sutton et al., [2021](#page-14-23)), in particular for competitive interactions (Hughes, [2012](#page-12-32)).

Although we limited our simulation to the juvenile stage, this does not imply that later stages can be neglected. Changing body size can have carry-over effects onto the next generation. For example, Wootton et al. ([2022](#page-14-24)) showed that smaller-sized female zebrafish under warming displayed larger gonads than same-sized females

 SASAKI ET AL. *PHONE CONTRACTE ALL CONTRACTE ACCOUNT* \mathbf{M}/\mathbf{H} **CONTRACTE ACCOUNT** \mathbf{M}/\mathbf{H} **CONTRACTE ACCOUNT** \mathbf{M}/\mathbf{H} **CONTRACTE ACCOUNT** \mathbf{M}/\mathbf{H} **CONTRACTE ACCOUNT** \mathbf{M}/\mathbf{H} **C**

at control temperatures, thus showing the potential for increased reproductive output under elevated temperatures. Conversely, smaller adult body sizes could also affect reproductive output negatively. For example, a smaller-sized mature female fish has been shown to reproduce disproportionally less than a larger female (Barneche et al., [2018](#page-11-16)). As such, it is important to assess the effect of changing body size on reproductive output and simulate the consequence of changing body size across generations.

We note that faster juvenile growth forecast in the current model may still lead to reduced maximum body size at later life stage (i.e., temperature-size rule, Atkinson, [1995](#page-11-17)). Higher but not stressful temperature has been found to shrink maximum body size in natural fish populations more strongly in active fish than sedentary fish (van Rijn et al., [2017\)](#page-14-25). Furthermore, Ikpewe et al. ([2021](#page-12-33)) found that increasing ocean temperatures enhanced length of juveniles of commercial species but decreased that of adults. As the adult stage was excluded from the current model, our findings of larger juvenile lengths for both tropical and temperate species should not directly be linked to adult size.

If the negative effects of novel interactions result in replacing native temperate species, this may lead to functional homogenisation where native species are replaced by non-native species (Olden et al., [2004](#page-13-28)). For example, our temperate species *Atypichthys strigatus*, is known as a generalist species as well as a facultative cleaner to other reef fish species (Glasby & Kingsford, [1994](#page-12-17)). Cleaner species provide important functions in the communities, such as enhancing the body condition of fish species (Grutter, [2010](#page-12-34); Ros et al., [2020](#page-13-29)) and increasing fish abundance (Bshary, [2003](#page-11-18)). Replacement of ecological specialist species by widespread generalist can modify the functional diversity of a community (Olden et al., [2004](#page-13-28)), and a decrease in functional diversity might diminish overall community and ecosystem functioning (Tilman et al., [1997](#page-14-26)), stability (Sankaran & McNaughton, [1999](#page-14-27)) and resistance to environmental change by simply narrowing the available range of species-specific response (Stachowicz et al., [2002](#page-14-28)). Although the warming-facilitated arrival of range-shifting species can increase species richness in the receiving communities (Hiddink & Ter Hofstede, [2008](#page-12-35); Steinbauer et al., [2018](#page-14-29); Telwala et al., [2013](#page-14-30)), it is important to understand the nature of novel interactions and how this may shape functional diversity in the community.

4.4 | **Model performance**

We found that the accuracy of our model in predicting the effect of changing temperature on fish growth in the field depended on the model design (with or without species interactions). For example, the model without species interaction predicted maximum size of juvenile *A. vaigiensis* at subtropics and mild-temperate better than the model with the interaction, but the latter predicted fish size better at warmtemperate latitudes than at subtropics and mild-temperate (Figure [S4](#page-15-0)). Given that our models already integrated the variation in thermal sensitivity (TA) and feeding rate (*f*), factors other than temperature

and the interaction between these two species likely generated the difference between the predicted and the observed maximum body lengths. Indeed, earlier work using the DEB model emphasized that elucidating the factors that are responsible for generating variation in organismal physiological processes is critical for predictions (Alunno-Bruscia et al., [2011](#page-11-19); Pecquerie et al., [2009](#page-13-30); Stubbs et al., [2020;](#page-14-31) Thomas & Bacher, [2018](#page-14-9)). For example, Pecquerie et al. ([2009](#page-13-30)) found that simulating the differences in hatching dates allowed them to obtain large observed variation in length at age 1 of anchovy. In our case, we selected the first of January for the recruitment to begin and initiated the simulation from the selected date consistently across all sampling site for both species. We used a single date instead of multiple dates because of limited understanding regarding the factors causing variation in the timing of recruitment in our species. Although it was possible for us to perform simulations with various recruitment start dates, uncertainty would likely propagate throughout the simulation and cause further difference between observed and predicted sizes, as previously suggested (Ren et al., [2020](#page-13-31)).

ogeography

Secondly, our model used standard starvation mode where starved individual use energy for development to pay for somatic maintenance until reserve energy falls below threshold. This starvation mode may differ from realistic energy use because fish are known to have developed alternative strategies for energy allocation under stressful conditions. For example, during prolonged energy deprivation, fish can divert energy from fat storage (short-term starvation) and muscle protein (long-term starvation) (Bar, [2014](#page-11-20); Furne & Sanz, [2017](#page-12-36)). These strategies mean that fish can survive even after losing muscle mass. Unfortunately, the starvation mode in our model did not allow diverting energy from the structural volume to sustain body function. Consequently, simulated individuals died soon after its alternative energy resource, maturity, was exhausted. Our model will greatly improve if we add new parameters accounting for the species-specific changes in energy requirement during starvation (e.g., Monaco et al., [2014](#page-13-32)). Implementing species-specific starvation mode may clarify the modulating effect of food intake, thus species interaction, on species response to temperature stress, as previously shown (Almeida et al., [2022](#page-11-21); Biro et al., [2021](#page-11-22); Hedström et al., [2017](#page-12-37); Shoup & Wahl, [2011](#page-14-32)).

Although we could not provide a one-size-fits-all model for predicting in situ growth of the tropical and the temperate species, this does not prevent our models from becoming a useful tool for predicting species performance in the field, as one of our two models still predicted in situ body sizes with a relatively low error rate. We suggest that, rather than relying on a single model, using two models and obtaining a range of body sizes will enhance the ability of our models to predict species response to environmental change that is closely matched with field observations.

4.5 | **Conclusions**

While the presence of tropical vagrant species in cool temperate regions represent a precursor to poleward range-extension, vagrants

are often excluded from predictive models as they are considered as noise. However, because these individuals interact and affect feeding rates of local species, excluding them from the models can neglect the important role of novel species interaction on species growth and population persistence under changing temperatures. We showed that using a dynamic energy budget approach, we can model the responses of tropical and interacting temperate species to future warming and novel interactions that are anticipated to occur in temperate coastal regions. Our study also highlighted the strong asset of the DEB model by showing its ability to simulate changes in fitness-related traits, as opposed to changes in habitat suitability as is the case for correlative species distribution models. Considering the strong link between body size and temperature, our study demonstrated a predictive model that provides a useful platform to simulate important fitness-related traits changes under changing environmental conditions.

ACKNOWLEDGEMENTS

This research was funded by an Australian Research Council (ARC) Discovery Project (DP170101722 and DP230101932) to IN and DJB. Fish were collected under NSW DPI permit F94/696, and experiments were performed under University of Technology Sydney Animal Ethics Application 17-117 and the University of Adelaide Animal Ethics Application # S-2018-107. We thank Angus Mitchell, Andrew Niccum, Sergio Torres Gabarda, Nigel Coombes and Amanda Scholes for support in running the aquarium experiments at the Sydney Institute of Marine Science. Open access publishing facilitated by The University of Adelaide, as part of the Wiley - The University of Adelaide agreement via the Council of Australian University Librarians.

CONFLICT OF INTEREST STATEMENT

We have no conflicts of interest to disclose.

DATA AVAILABILITY STATEMENT

Parameters used for simulation is available in Table [S2](#page-15-0).

ORCID

Ivan Nagelkerken <https://orcid.org/0000-0003-4499-3940>

REFERENCES

- Alexander, J. M., Diez, J. M., & Levine, J. M. (2015). Novel competitors shape species' responses to climate change. *Nature*, *525*, 515–518.
- Allen, G. R., & Erdmann, M. V. (2012). *Reef fishes of the East Indies*. University of Hawai'i Press, Volumes I-III. Tropical Reef Research.
- Almeida, J., Lopes, A. R., Ribeiro, L., Castanho, S., Candeias-Mendes, A., Pousão-Ferreira, P., & Faria, A. M. (2022). Effects of exposure to elevated temperature and different food levels on the escape response and metabolism of early life stages of white seabream, *Diplodus sargus*. *Conservation Physiology*, *10*(1), coac023.
- Alunno-Bruscia, M., Van der Veer, H. W., & Kooijman, S. A. (2011). The AquaDEB project: Physiological flexibility of aquatic animals analysed with a generic dynamic energy budget model (phase II). *Journal of Sea Research*, *66*(4), 263–269.
- Araújo, M. B., & Luoto, M. (2007). The importance of biotic interactions for modelling species distributions under climate change. *Global Ecology and Biogeography*, *16*(6), 743–753.
- Araújo, M. B., & Pearson, R. G. (2005). Equilibrium of species' distributions with climate. *Ecography*, *28*, 693–695.
- Araújo, M. B., Pearson, R. G., Thuiller, W., & Erhard, M. (2005). Validation of species–climate impact models under climate change. *Global Change Biology*, *11*, 1504–1513.
- Aryal, A., Shrestha, U. B., Ji, W., Ale, S. B., Shrestha, S., Ingty, T., Maraseni, T., Cockfield, G., & Raubenheimer, D. (2016). Predicting the distributions of predator (snow leopard) and prey (blue sheep) under climate change in the Himalaya. *Ecology and Evolution*, *6*(12), 4065– 4075.<https://doi.org/10.1002/ece3.2196>
- Atkinson, D. (1995). Effects of temperature on the size of aquatic ectotherms: Exceptions to the general rule. *Journal of Thermal Biology*, *20*(1/2), 61–74.
- Atlas of Living Australia. (2021a). Atlas of Living Australia occurrence. [https://biocache.ala.org.au/occurrences/search?q](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3A268b9721-ffc9-41f1-a58c-f30fde3241dd&qualityProfile=ALA)=lsid%3Aurn% [3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3A268b9721-ffc9-](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3A268b9721-ffc9-41f1-a58c-f30fde3241dd&qualityProfile=ALA) [41f1-a58c-f30fde3241dd&qualityProfile](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3A268b9721-ffc9-41f1-a58c-f30fde3241dd&qualityProfile=ALA)=ALA. Accessed 29 December 2021.
- Atlas of Living Australia. (2021b). Atlas of Living Australia occurrence. [https://biocache.ala.org.au/occurrences/search?q](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3Ab810d4f6-b879-4c48-94f2-ba8756290100&qualityProfile=ALA)=lsid%3Aurn% [3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3Ab810d4f6-b879-](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3Ab810d4f6-b879-4c48-94f2-ba8756290100&qualityProfile=ALA) [4c48-94f2-ba8756290100&qualityProfile](https://biocache.ala.org.au/occurrences/search?q=lsid%3Aurn%3Alsid%3Abiodiversity.org.au%3Aafd.taxon%3Ab810d4f6-b879-4c48-94f2-ba8756290100&qualityProfile=ALA)=ALA. Accessed 29 December 2021.
- Bar, N. (2014). Physiological and hormonal changes during prolonged starvation in fish. *Canadian Journal of Fisheries and Aquatic Sciences*, *71*(10), 1447–1458.
- Barneche, D. R., Robertson, D. R., White, C. R., & Marshall, D. J. (2018). Fish reproductive-energy output increases disproportionately with body size. *Science*, *360*(6389), 642–645.
- Bastille-Rousseau, G., Schaefer, J. A., Peers, M. J. L., Ellington, E. H., Mumma, M. A., Rayl, N. D., Mahoney, S. P., & Murray, D. L. (2018). Climate change can alter predator-prey dynamics and population viability of prey. *Oecologia*, *186*, 141–150.
- Beale, C. M., & Lennon, J. J. (2012). Incorporating uncertainty in predictive species distribution modelling. *Philosophical Transactions of the Royal Society B*, *367*, 247–258. [https://doi.org/10.1098/rstb.2011.](https://doi.org/10.1098/rstb.2011.0178) [0178](https://doi.org/10.1098/rstb.2011.0178)
- Biro, P. A., Post, J. R., & Beckmann, C. (2021). Autumn lipid reserves, overwinter lipid depletion, and high winter mortality of rainbow trout in experimental lakes. *Canadian Journal of Fisheries and Aquatic Sciences*, *78*(6), 738–743.
- Booth, D. J., Beretta, A., Brown, L., & Figueira, W. F. (2018). Predicting success of range-expanding coral reef fish in temperature habitat using temperature-abundance relationships. *Frontiers in Marine Science*, *5*, 31.<https://doi.org/10.3389/fmars.2018.00031>
- Bray, D., Gomon, M., (2011). *Atypichthys strigatus* Mado in Museums Victoria Collections, [https://collections.museumsvictoria.com.au/](https://collections.museumsvictoria.com.au/species/8319) [species/8319.](https://collections.museumsvictoria.com.au/species/8319) Accessed 08 November 2022.
- Bray, D.J. (2023). *Atypichthys strigatus* in fishes of Australia. Accessed 04 Jun 2024, [https://fishesofaustralia.net.au/home/](https://fishesofaustralia.net.au/home/species/467) [species/467](https://fishesofaustralia.net.au/home/species/467)
- Bshary, R. (2003). The cleaner wrasse, *Labroides dimidiatus*, is a key organism for reef fish diversity at Ras Mohammed National Park, Egypt. *Journal of Animal Ecology*, *72*(1), 169–176.
- Caplat, P., Anand, M., & Bauch, C. (2008). Interactions between climate change, competition, dispersal, and disturbances in a tree migration model. *Theoretical Ecology*, *1*, 209–220.
- Chen, I. C., Hill, J. K., Ohlemüller, R., Roy, D. B., & Thomas, C. D. (2011). Rapid range shifts of species associated with high levels of climate warming. *Science*, *333*, 1024–1026.
- Chivers, W., Walne, A., & Hays, G. (2017). Mismatch between marine plankton range movements and the velocity of climate change. *Nature Communications*, *8*, 14434.
- Citores, L., Ibaibarriaga, L., Lee, D. J., Brewer, M. J., Santos, M., & Chust, G. (2020). Modelling species presence–absence in the ecological niche theory framework using shape-constrained generalized additive models. *Ecological Modelling*, *418*, 108926.
- Coni, E. O. C., Booth, D. J., & Nagelkerken, I. (2021). Novel species interactions and environmental conditions reduce foraging competency at the temperate range edge of a range-extending coral reef fish. *Coral Reefs*, *40*(5), 1525–1536.
- Cooper, J. C., & Soberón, J. (2017). Creating individual accessible area hypotheses improves staked species distribution model performance. *Global Ecology and Biogeography*, *27*(1), 156–165.
- Cyr, H. (2000). Individual energy use and the allometry of population density. In *Scaling in biology* (pp. 267–295). Oxford University Press, Inc.
- Cyr, H., Peters, R. H., & Downing, J. A. (1997). Population density and community size structure: Comparison of aquatic and terrestrial systems. *Oikos*, *80*, 139–149.
- D'Amen, M., & Azzurro, E. (2019). Integrating univariate niche dynamics in species distribution models: A step forward for marine research on biological invasions. *Journal of Biogeography*, *47*(3), 686–697.
- de Souza, J. S., & Dos Santos, L. N. (2023). Resident species, not immigrants, drive reorganization of estuarine fish assemblages in response to warming. *Ecology*, *104*(5), e3987.
- Dolci, D., & Peruzzi, L. (2022). Assessing the effectiveness of correlative ecological niche model temporal projection through floristic data. *Biology*, *11*(8), 1219.
- Donelson, J. M., Sunday, J. M., Figueira, W. F., Gaitán-Espitia, J. D., Hobday, A. J., Johnson, C. R., Leis, J. M., Ling, S. D., Marshall, D., Pandolfi, J. M., Pecl, G., Rodgers, G. G., Booth, D. J., & Munday, P. L. (2019). Understanding interactions between plasticity, adaptation and range shifts in response to marine environmental change. *Philosophical Transactions of the Royal Society B*, *374*(1768), 20180186.
- Donelson, J. M., Munday, P. L., McCormick, M. I., & Nilsson, G. E. (2011). Acclimation to predicted ocean warming through developmental plasticity in a tropical reef fish. *Global Change Biology*, *17*(4), 1712–1719.
- Figueira, W. F., Biro, P., Booth, D. J., & Valenzuela, V. C. (2009). Performance of tropical fish recruiting to temperate habitats: Role of ambient temperature and implications of climate change. *Marine Ecology Progress Series*, *384*, 231–239.
- Figueira, W. F., & Booth, D. J. (2010). Increasing ocean temperatures allow tropical fishes to survive overwinter in temperate waters. *Global Change Biology*, *16*, 506–516.
- Freeman, B. G., Lee-Yaw, J. A., Sunday, J. M., & Hargreaves, A. L. (2018). Expanding, shifting and shrinking: The impact of global warming on species' elevational distributions. *Global Ecology and Biogeography*, *27*, 1268–1276.<https://doi.org/10.1111/geb.12774>
- Froese, R., & Pauly, D. (2000). *FishBase 2000: concepts, design and data sources* (p. 344). ICLARM.
- Furne, M., & Sanz, A. (2017). Starvation in fish – Sturgeon and rainbow trout as examples. In V. Preedy & V. Patel (Eds.), *Handbook of ramine, starvation, and nutrient deprivation*. Springer.
- Gervais, C. R., Champion, C., & Pecl, G. T. (2021). Species on the move around the Australian coastline: A continental-scale review of climate-driven species redistribution in marine systems. *Global Change Biology*, *27*(14), 3200–3217.
- Gillard, M., Thiébaut, G., Deleu, C., & Leroy, B. (2017). Present and future distribution of three aquatic plants taxa across the world: Decrease in native and increase in invasive ranges. *Biological Invasions*, *19*, 2159–2170.
- Glasby, T. M., & Kingsford, M. J. (1994). *Atypichthys strigatus* (Pisces: Scorpididae): An opportunistic planktivore that responds to benthic disturbances and cleans other fishes. *Australian Journal of Ecology*, *19*(4), 385–394.

Grutter, A. S. (2010). Cleaner fish. *Current Biology*, *20*(13), R547–R549.

ogeography

- Guisan, A., Lehmann, A., Ferrier, S., Austin, M., Overton, J. M. C., Aspinall, R., & Hastie, T. (2006). Making better biogeographical predictions of species' distributions. *Journal of Applied Ecology*, *43*(3), 386–392.
- Guisan, A., Petitpierre, B., Broennimann, O., Daehler, C., & Kueffer, C. (2014). Unifying niche shift studies: Insights from biological invasions. *Trends in Ecology & Evolution*, *29*(5), 260–269.
- Guisan, A., & Zimmermann, N. E. (2000). Predictive habitat distribution models in ecology. *Ecological Modelling*, *135*, 147–186.
- Hedström, P., Bystedt, D., Karlsson, J., Bokma, F., & Byström, P. (2017). Brownification increases winter mortality in fish. *Oecologia*, *183*(2), 587–595.
- Hiddink, J. G., & Ter Hofstede, R. (2008). Climate induced increases in species richness of marine fishes. *Global Change Biology*, *14*(3), 453–460.
- Holbrook, S. J., & Schmitt, R. J. (1989). Resource overlap, prey dynamics, and the strength of competition. *Ecology*, *70*, 1943–1953.
- Hollebone, A. L., & Hay, M. E. (2008). An invasive crab alters interaction webs in a marine community. *Biological Invasions*, *10*, 347–358.
- Hughes, L. (2012). Climate change impacts on species interactions: Assessing the threat of cascading extinctions. In L. Hannah (Ed.), *Saving a Million Species* (pp. 337–359). Island Press/Center for Resource Economics.
- Hyndes, G. A., Heck, K. L., Vergés, A., Harvey, E. S., Kendrick, G. A., Lavery, P. S., McMahon, K., Orth, R. J., Pearce, A., Vanderklift, M., Wernberg, T., Whiting, S., & Wilson, S. (2016). Accelerating tropicalization and the transformation of temperate seagrass meadows. *Bioscience*, *66*(11), 938–948.
- Ikpewe, I. E., Baudron, A. R., Ponchon, A., & Fernandes, P. G. (2021). Bigger juveniles and smaller adults: Changes in fish size correlate with warming seas. *Journal of Applied Ecology*, *58*(4), 847–856.
- IMOS. (2019). SRS – SST – L3S – Single Sensor – 6 day – day and night time – Australia. <https://portal.aodn.org.au/search>. Accessed 18 Dec 2019.
- Johnson, C. R., Banks, S. C., Barrett, N. S., Cazassus, F., Dunstan, P. K., Edgar, G. J., Frusher, S. D., Gardner, C., Haddon, M., Helidoniotis, F., & Hill, K. L. (2011). Climate change cascades: Shifts in oceanography, species' ranges and subtidal marine community dynamics in eastern Tasmania. *Journal of Experimental Marine Biology and Ecology*, *400*(1–2), 17–32.
- Kearney, M., Phillips, B. L., Tracy, C. R., Christian, K. A., Betts, G., & Porter, W. P. (2008). Modelling species distributions without using species distributions: The cane toad in Australia under current and future climates. *Ecography*, *31*(4), 423–434.
- Kingsbury, K. M., Gillanders, B. M., Booth, D. J., Coni, E. O. C., & Nagelkerken, I. (2020). Range-extending coral reef fishes trade-off growth for maintenance of body condition in cooler waters. *Science of the Total Environment*, *703*, 134598.
- Kingsbury, K. M., Gillanders, B. M., Booth, D. J., & Nagelkerken, I. (2019). Trophic niche segregation allows range-extending coral reef fishes to co-exit with temperate species under climate change. *Global Change Biology*, *26*(2), 721–733.
- Kohda, M., Shibata, J., Awata, S., Gomagano, D., Takeyama, T., Hori, M., & Heg, D. (2008). Niche differentiation depends on body size in a cichlid fish: A model system of a community structured according to size regularities. *Journal of Animal Ecology*, *77*, 859–868.
- Kooijman, S. A. L. M. (2010). *Dynamic energy budget theory for metabolic organisation* (3rd ed.). Cambridge University Press.
- Kordas, R. L., Harley, C. D. G., & O'Connor, M. I. (2011). Community ecology in a warming world: The influence of temperature on interspecific interactions in marine systems. *Journal of Experimental Marine Biology and Ecology*, *400*, 218–226.
- Lee-Yaw, A. J., McCune, L. J., Pironon, S., & Sheth, N. S. (2022). Species distribution models rarely predict the biology of real populations. *Ecography*, *2022*(6), e05877.

 burnal of the contract of th

2256 [|] SASAKI et al.

- Liang, Y., Duveneck, M. J., Gustafson, E. J., Serra-Diaz, J. M., & Thompson, J. R. (2017). How disturbance, competition, and dispersal interact to prevent tree range boundaries from keeping pace with climate change. *Global Change Biology*, *24*(1), e335–e351.
- Ling, S. D. (2008). Range expansion of a habitat-modifying species leads to loss of taxonomic diversity: A new and impoverished reef state. *Oecologia*, *156*, 883–894.
- Liu, C., Wolter, C., Xian, W., & Jeschke, J. M. (2020). Species distribution models have limited spatial transferability for invasive species. *Ecology Letters*, *23*(11), 1682–1692.
- Manes, S., Costello, M. J., Beckett, H., Debnath, A., Devenish-Nelson, E., Grey, K. A., Jenkins, R., Khan, T. M., Kiessling, W., Krause, C., & Maharaj, S. S. (2021). Endemism increases species' climate change risk in areas of global biodiversity importance. *Biological Conservation*, *257*, 109070.
- Marques, G., Augustine, S., Lika, K., Pecquerie, L., Dominogs, T., & Kooijman, S. A. L. M. (2018). The AmP project: Comparing species on the basis of dynamic energy budget parameters. *PLoS Computational Biology*, *14*(5), e1006100.
- Martin, B., Zimmer, E., Grimm, V., & Jager, T. (2010). DEB-IBM user manual. [https://cream-itn.eu/creamwp/wp-content/uploads/DEB-](https://cream-itn.eu/creamwp/wp-content/uploads/DEB-IBM-user-manual.pdf)[IBM-user-manual.pdf](https://cream-itn.eu/creamwp/wp-content/uploads/DEB-IBM-user-manual.pdf)
- Martin, B. T., Zimmer, E. I., Grimm, V., & Jager, T. (2012). Dynamic Energy Budget theory meets individual-based modelling: A generic and accessible implementation. *Methods in Ecology and Evolution*, *3*, 445–449.
- Maxwell, S. L., Butt, N., Maron, M., McAlpine, C. A., Chapman, S., Ullmann, A., Segan, D. B., & Watson, J. E. (2019). Conservation implications of ecological responses to extreme weather and climate events. *Diversity and Distributions*, *25*(4), 613–625.
- Monaco, C. J., Booth, D. J., Figueira, W. F., Gillanders, B. M., Schoeman, D. S., Bradshaw, C., & Nagelkerken, I. (2020). Natural and anthropogenic climate variability shape assemblages of range-extending coral-reef fishes. *Journal of Biogeography*, *48*, 1063–1075.
- Monaco, C. J., & McQuaid, C. D. (2019). Climate warming reduces the reproductive advantage of a globally invasive intertidal mussel. *Biological Invasions*, *21*(7), 2503–2516.
- Monaco, C. J., Wethey, D. S., & Helmuth, B. (2014). A dynamic energy budget (DEB) model for the keystone predator *Pisaster ochraceus*. *PLoS One*, *9*(8), e104658.
- Monaco, C. J., Wethey, D. S., & Helmuth, B. (2016). Thermal sensitivity and the role of behaviour in driving an intertidal predator-prey interaction. *Ecological Monographs*, *86*(4), 429–447.
- Moullec, F., Barrier, N., Drira, S., Guilhaumon, F., Marsaleix, P., Somot, S., Ulses, C., Velez, L., & Shin, Y. J. (2019). An end-to-end model reveals losers and winners in a warming Mediterranean Sea. *Frontiers in Marine Science*, *6*, 345. [https://doi.org/10.3389/fmars.](https://doi.org/10.3389/fmars.2019.00345) [2019.00345](https://doi.org/10.3389/fmars.2019.00345)
- Neuheimer, A. B., Thresher, R. E., Lyle, J. M., & Semmens, J. M. (2011). Tolerance limit for fish growth exceeded by warming waters. *Nature Climate Change*, *1*(2), 110–113.
- Nye, J. A., Link, J. S., Hare, J. A., & Overholtz, W. J. (2009). Changing spatial distribution of fish stocks in relation to climate and population size on the Northeast United States continental shelf. *Marine Ecology Progress Series*, *393*, 111–129.
- Olden, J. D., Poff, N. L., Douglas, M. R., Douglas, M. E., & Fausch, K. D. (2004). Ecological and evolutionary consequences of biotic homogenization. *Trends in Ecology & Evolution*, *19*(1), 18–24.
- Osland, M. J., Stevens, P. W., Lamont, M. M., Brusca, R. C., Hart, K. M., Waddle, J. H., Langtimm, C. A., Williams, C. M., Keim, B. D., Terando, A. J., & Reyier, E. A. (2021). Tropicalization of temperate ecosystems in North America: The northward range expansion of tropical organisms in response to warming winter temperatures. *Global Change Biology*, *27*(13), 3009–3034.
- Pörtner, H. O., & Knust, R. (2007). Climate change affects marine fishes through the oxygen limitation of thermal tolerance. *Science*, *315*(5808), 95–97.
- Parmesan, C. (2006). Ecological and evolutionary responses to recent climate change. *Annual Review of Ecology, Evolution, and Systematics*, *37*, 637–669.
- Payne, N. L., Smith, J. A., van der Meulen, D. E., Taylor, M. D., Watanabe, Y. Y., Takahashi, A., Marzullo, T. A., Gray, C. A., Cadiou, G., & Suthers, I. M. (2016). Temperature dependence of fish performance in the wild: Links with species biogeography and physiological thermal tolerance. *Functional Ecology*, *30*(6), 903–912.
- Pearce, A., Hutchins, B., Hoschke, A., & Fearns, P. (2016). Record high damselfish recruitment at Rottnest Island, Western Australia, and the potential for climate-induced range extension. *Regional Studies in Marine Science*, *8*, 77–88.
- Pecquerie, L., Petitgas, P., & Kooijman, S. A. L. M. (2009). Modeling fish growth and reproduction in the context of the Dynamic Energy Budget theory to predict environmental impact on anchovy spawning duration. *Journal of Sea Research*, *62*(2–3), 93–105.
- Perry, A. L., Low, P. J., Ellis, J. R., & Reynolds, J. D. (2005). Climate change and distribution shifts in marine fishes. *Science*, *308*(5730), 1912–1915.
- Peterson, A. T., & Vieglais, D. A. (2001). Predicting species invasions using ecological niche modeling: New approaches from bioinformatics attack a pressing problem: A new approach to ecological niche modelling, based on new tools drawn from biodiversity informatics, is applied to the challenge of predicting potential species' invasions. *Bioscience*, *51*, 363–371.
- Pinsky, M. L., Selden, R. L., & Kitchel, Z. J. (2020). Climate-driven shifts in marine species ranges: Scaling from organisms to communities. *Annual Review of Marine Science*, *12*, 153–179.
- Pinsky, M. L., Worm, B., Fogarty, M. J., Sarmiento, J. L., & Levin, S. A. (2013). Marine taxa track local climate velocities. *Science*, *341*(6151), 1239–1242.
- Poloczanska, E. S., Brown, C. J., Sydeman, W. J., Kiessling, W., Shoeman, D. S., Moore, P. J., Brander, K., Bruno, J. F., Buckley, L. B., Burrows, M. T., Duarte, C. M., Halpern, B. S., Holding, J., Kappel, C. V., O'Connor, M. I., Pandolfi, J. M., Parmesan, C., Schwing, F., Thompson, S. A., & Richardson, A. (2013). Global imprint of climate change on marine life. *Nature Climate Change*, *3*, 919–925. [https://doi.org/10.1038/](https://doi.org/10.1038/NCLIMATE1958) [NCLIMATE1958](https://doi.org/10.1038/NCLIMATE1958)
- Ren, J. S., Jin, X., Yang, T., Kooijman, S. A. L. M., & Shan, X. (2020). A dynamic energy budget model for small yellow croaker *Larimichthys polyactis*: Parameterisation and application in its main geographic distribution waters. *Ecological Modelling*, *427*, 109051.
- Ridgway, K. R. (2007). Long-term trend and decadal variability of the southward penetration of the East Australia current. *Geophysical Research Letters*, *34*, L13613.
- Ros, A. F., Nusbaumer, D., Triki, Z., Grutter, A. S., & Bshary, R. (2020). The impact of long-term reduced access to cleaner fish on health indicators of resident client fish. *Journal of Experimental Biology*, *223*(24), ieb231613.
- Rose, P. M., Kennard, M. J., Moffatt, D. B., Sheldon, F., & Bulter, G. L. (2016). Testing three species distribution modelling strategies to define fish assemblage reference conditions for stream bioassessment and related applications. *PLoS One*, *11*(1), e0146728.
- Sánchez-Hernández, J., Gabler, H. M., & Amundsen, P. A. (2017). Prey diversity as a driver of resource partitioning between river-dwelling fish species. *Ecology and Evolution*, *7*, 2058–2068.
- Sanches, F. H. C., Miyai, C. A., Coasta, T. M., Christofoletti, R. A., Volpato, G. L., & Barreto, R. E. (2012). Aggressiveness overcomes body-size effects in fights staged between invasive and native fish species with overlapping niches. *PLoS One*, *7*(1), e29746. [https://doi.org/10.](https://doi.org/10.1371/journal.pone.0029746) [1371/journal.pone.0029746](https://doi.org/10.1371/journal.pone.0029746)

articles are governed by the applicable Creative Commons

- Sandblom, E., Clark, T. D., Gräns, A., Ekström, A., Brijs, J., Sundström, L. F., Odelström, A., Adill, A., Aho, T., & Jutfelt, F. (2016). Physiological constraints to climate warming in fish follow principles of plastic floors and concrete ceilings. *Nature Communications*, *7*(1), 11447.
- Sankaran, M., & McNaughton, S. J. (1999). Determinants of biodiversity regulate compositional stability of communities. *Nature*, *401*, 691–693.
- Sasaki, M., Kingsbury, K. M., Booth, D. J., & Nagelkerken, I. (2024). Body size mediates trophic interaction strength of novel fish assemblages under climate change. *Journal of Animal Ecology*, *93*, 705–714.
- Seneviratne, S. I., Zhang, X., Adnan, M., Badi, W., Dereczynski, C., Di Luca, A., Ghosh, S., Iskandar, I., Kossin, J., Lewis, S., Otto, F., Pinto, I., Satoh, M., Vicente-Serrano, S. M., Wehner, M., & Zhou, B. (2021). Weather and climate extreme events in a changing climate. In V. Masson-Delmotte, P. Zhai, A. Pirani, S. L. Connors, C. Péan, S. Berger, N. Caud, Y. Chen, L. Goldfarb, M. I. Gomis, M. Huang, K. Leitzell, E. Lonnoy, J. B. R. Matthews, T. K. Maycock, T. Waterfield, O. Yelekçi, R. Yu, & B. Zhou (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* (pp. 1513– 1766). Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA.
- Shoup, D. E., & Wahl, D. H. (2011). Body size, food, and temperature affect overwinter survival of age-0 bluegills. *Transactions of the American Fisheries Society*, *140*(5), 1298–1304.
- Sinclair, B. J., Marshall, K. E., Sewell, M. A., Levesque, D. L., Willett, C. S., Slotsbo, S., Dong, Y., Harley, C. D., Marshall, D. J., Helmuth, B. S., & Huey, R. B. (2016). Can we predict ectotherm responses to climate change using thermal performance curves and body temperatures? *Ecology Letters*, *19*(11), 1372–1385.
- Smale, D. A., & Wernberg, T. (2013). Extreme climatic event drives range contraction of a habitat-forming species. *Proceedings of the Royal Society B: Biological Sciences*, *280*(1754), 20122829.
- Smith, S. M., Fox, R. J., Booth, D. J., & Donelson, J. M. (2017). Stick with your own kind, or hand with the locals? Implications of shoaling strategy for tropical reef fish on a range-expansion frontline. *Global Change Biology*, *24*(4), 1663–1672.
- Stachowicz, J. J., Fried, H., Osman, R. W., & Whitlatch, R. B. (2002). Biodiversity, invasion resistance, and marine ecosystem function: Reconciling pattern and process. *Ecology*, *83*(9), 2575–2590.
- Steinbauer, M. J., Grytnes, J. A., Jurasinski, G., Kulonen, A., Lenoir, J., Pauli, H., Rixen, C., Winkler, M., Bardy-Durchhalter, M., Barni, E., & Bjorkman, A. D. (2018). Accelerated increase in plant species richness on mountain summits is linked to warming. *Nature*, *556*(7700), 231–234.
- Stewart, E. M., Frasca, V. R., Wilson, C. C., & Raby, G. D. (2023). Shortterm acclimation dynamics in a coldwater fish. *Journal of Thermal Biology*, *112*, 103482.
- Stubbs, J. L., Marn, N., Vanderklift, M. A., Fossette, S., & Mitchell, N. J. (2020). Simulated growth and reproduction of green turtles (*Chelonia mydas*) under climate change and marine heatwave scenarios. *Ecological Modelling*, *431*, 109185.
- Sutton, A. O., Studd, E. K., Fernandes, T., Bates, A. E., Bramburger, A. J., Cooke, S. J., Hayden, B., Henry, H. A. L., Humphries, M. M., Martin, R., McMeans, B., Moise, E., O'Sullivan, A. M., Sharma, S., & Templer, P. H. (2021). Frozen out: Unanswered questions about winter biology. *Environmental Reviews*, *29*(4), 431–442.
- Telwala, Y., Brook, B. W., Manish, K., & Pandit, M. K. (2013). Climateinduced elevational range shifts and increase in plant species richness in a Himalayan biodiversity epicentre. *PLoS One*, *8*(2), e57103.
- Thomas, Y., & Bacher, C. (2018). Assessing the sensitivity of bivalve populations to global warming using an individual-based modelling approach. *Global Change Biology*, *24*(10), 4581–4597.

Tilman, D., Knops, J., Wedin, D., Reich, P., Ritchie, M., & Siemann, E. (1997). The influence of functional diversity and composition on ecosystem processes. *Science*, *277*(5330), 1300–1302.

ogeography

- van Rijn, I., Buba, Y., DeLong, J., Kiflawi, M., & Belmaker, J. (2017). Large but uneven reduction in fish size across species in relation to changing sea temperatures. *Global Change Biology*, *23*(9), 3667–3674.
- Van der Meer, J. (2006). An introduction to dynamic energy budget (DEB) models with special emphasis on parameter estimation. *Journal of Sea Research*, *56*(2), 85–102.
- Van Der Walt, K. A., Potts, W. M., Porri, F., Winkler, A. C., Duncan, M. I., Skeeles, M. R., & James, N. C. (2021). Marine heatwaves exceed cardiac thermal limits of adult Sparid fish (*Diplodus capensis*, Smith 1884). *Frontiers in Marine Science*, *8*, 801.
- Vergés, A., Stenberg, P. D., Hay, M. E., Poore, A. G. B., Campbell, A. H., Ballesteros, E., Heck, K. L., Jr., Booth, D. J., Coleman, M. A., Feary, D. A., Figueira, W., Langlois, T., Marzinelli, E. M., Mizerek, T., Mumby, P. J., Nakamura, Y., Roughan, M., van Sebille, E., Gupta, A. S., … Wilson, S. K. (2014). The tropicalization of temperate marine ecosystems: Climate-mediated changes in herbivory and community phase shifts. *Proceedings of the Royal Society of London, Series B: Biological Sciences*, *281*(1789), 20140846.
- Volkoff, H., & Rønnestad, I. (2020). Effects of temperature on feeding and digestive processes in fish. *Temperature*, *7*(4), 307–320. [https://](https://doi.org/10.1080/23328940.2020.1765950) doi.org/10.1080/23328940.2020.1765950
- WDC CLIMATE (2019). Public website. [https://cera-www.dkrz.de/](https://cera-www.dkrz.de/WDCC/ui/cerasearch/) [WDCC/ui/cerasearch/](https://cera-www.dkrz.de/WDCC/ui/cerasearch/). Accessed 18 Dec 2019.
- Wernberg, T., Smale, D. A., Tuya, F., Thomsen, M. S., Langlois, T. J., De Bettignies, T., Bennett, S., & Rousseaux, C. S. (2013). An extreme climatic event alters marine ecosystem structure in a global biodiversity hotspot. *Nature Climate Change*, *3*(1), 78–82.
- Whitfield, A. K., James, N. C., Lamberth, S. J., Adams, J. B., Perissinotto, R., Rajkaran, A., & Bornman, T. G. (2016). The role of pioneers as indicators of biogeographic range expansion caused by global change in southern African coastal waters. *Estuarine, Coastal and Shelf Science*, *172*, 138–153.
- Whitney, K. D., & Gabler, C. A. (2008). Rapid evolution in introduced species, 'invasive traits' and recipient communities: Challenges for predicting invasive potential. *Diversity and Distributions*, *14*(4), 569–580.
- Wilensky, U., (1999). NetLogo. Center for Connected Learning and Computer-Based Modeling, Northwestern University. [http://ccl.](http://ccl.northwestern.edu/netlogo/) [northwestern.edu/netlogo/](http://ccl.northwestern.edu/netlogo/).
- Williams, J. W., & Jackson, S. T. (2007). Novel climates, no-analog communities, and ecological surprises. *Frontiers in Ecology and the Environment*, *5*(9), 475–482.
- Wissinger, S. A. (1988). Life history and size structure of larval dragonfly populations. *Journal of the North American Benthological Society*, *7*(1), 13–28.
- Woodward, G., & Hildrew, A. G. (2002). Body-size determinants of niche overlap and intraguild predation within a complex food web. *Journal of Animal Ecology*, *71*, 1063–1074.
- Wootton, H. F., Morrongiello, J. R., Schmitt, T., & Audzijonyte, A. (2022). Smaller adult fish size in warmer water is not explained by elevated metabolism. *Ecology Letters*, *25*(5), 1177–1188.
- Worm, B., & Lotze, H. K. (2021). Marine biodiversity and climate change. In *Climate change* (pp. 445–464). Elsevier.
- Zeidberg, L. D., & Robison, B. H. (2007). Invasive range expansion by the Humboldt squid, *Dosidicus gigas*, in the eastern North Pacific. *Proceedings of the National Academy of Sciences of the United States of America*, *104*(31), 12948–12950.
- Zurell, D., Elith, J., & Schröder, B. (2012). Predicting to new environments: Tools for visualizing model behaviour and impacts on mapped distributions. *Diversity and Distributions*, *18*(6), 628–634.

BIOSKETCHES

Minami Sasaki is interested in marine ecology and the effect of environmental change on fish species. This work represents a component of her PhD work at the University of Adelaide on the effect of climate change and novel interactions on co-occurring range-shifting tropical fish species and resident temperate fish species.

Cristian Monaco is a physiological ecologist at the French National Institute of Ocean Science and Technology (Ifremer), working towards better understanding individual-level mechanisms that ultimately dictate organisms' responses to environmental forcings. By combining empirical data and mathematical models, Cristian is currently developing tools for predicting the impacts of climate change on tropical, marine aquaculture species.

David Booth is a fish ecologist at the University of Technology Sydney, interested in impacts of climate change on coral and temperate reef fishes and estuarine fishes and role of marine protected areas on fish conservation

Ivan Nagelkerken is a fish ecologist at the University of Adelaide, interested in the effects of climate change on fish communities and marine ecosystems. He also works on intertidal habitat connectivity for fishes.

Author contributions: Minami Sasaki, David J Booth and Ivan Nagelkerken conceived the ideas for the manuscript. Minami Sasaki analysed the data and simulated the DEB model with help from Cristián J Monaco. Minami Sasaki wrote the manuscript and all authors contributed critically to the drafts and gave final approval for publication.

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Sasaki, M., Monaco, C. J., Booth, D. J., & Nagelkerken, I. (2024). Ocean warming and novel species interactions boost growth and persistence of range-extending tropical fishes but challenge that of sympatric temperate species in temperate waters. *Journal of Biogeography*, *51*, 2243–2258. <https://doi.org/10.1111/jbi.14983>