RESEARCH PAPER

Ocean warming affects the distribution and abundance of resident fishes by changing their reproductive scope

Warren M. Potts · Anthony J. Booth · Timothy J. Richardson · Warwick H. H. Sauer

Received: 11 September 2012/Accepted: 7 October 2013/Published online: 15 October 2013 © Springer Science+Business Media Dordrecht 2013

Abstract With ocean warming predicted globally, one of the mechanisms driving distributional shifts and changes in the abundance of resident fishes is reproductive output. The relationship between sea surface temperature and the reproductive activity of a eurythermic, resident coastal species, blacktail seabream Diplodus sargus capensis, was examined in the "ocean warming" hotspot of the northern Benguela. Reproductive activity was found to be restricted to periods when the water temperature dropped below 20 °C. A metadata analysis conducted on the D. sargus sub-species complex similarly showed that reproductive activity was restricted to temperatures between 15 and 20 °C, regardless of the range in ambient water temperature. Based on these findings and using satellite derived SST information, we examined D. s. capensis's total and seasonal "reproductive scope" that is defined as either the area suitable for spawning each year or the duration of its potential spawning season at a fixed geographical locality, respectively. Trends were examined over the last three decades. Reproductive scope by area was found to be shrinking at a rate of 7 % per decade in southern Angola and expanding at a rate of 6 % per

W. M. Potts $(\boxtimes) \cdot A$. J. Booth $\cdot T$. J. Richardson \cdot

W. H. H. Sauer

Department of Ichthyology and Fisheries Science, Rhodes University, P.O. Box 94, Grahamstown 6140, South Africa e-mail: w.potts@ru.ac.za decade in northern Namibia. Reproductive scope by season decreased by 1.05 months per decade in Namibe, southern Angola and increased by 0.76 months per decade in Hentiesbaai, northern Namibia. Changes in reproductive scope may be a driving mechanism of distributional shifts in resident fishes, although the rate of the shifts is likely to be slow. More importantly, changes in reproductive scope will not be uniform throughout fish distributions and will most likely result in heterogeneous variations in fish abundance.

Keywords Angola · Blacktail seabream · Global climate change · Sea surface temperature

Introduction

Although it is accepted that climate change will have a major influence on the marine environment and its biota (Pörtner and Peck 2010), there is a paucity of predictive information on its potential impacts. In fishes, this development of predictive information is complicated by their diverse habitats, physiology and life histories. When coupled with the diverse regional climate variation, population- and area-specific research appears to be a prerequisite for the development of predictive models (Mcfarlane et al. 2000).

Of all of the climate driven physical and chemical stressors, changes in the ocean temperature is considered to be the most influential with a growing body of evidence highlighting its influence on fish distribution patterns (Stebbing et al. 2002; Perry et al. 2005; Last et al. 2011). This is hardly surprising as water temperature changes are known to affect several physiological processes in fishes, including their metabolic rate, fluidity of membranes and organ function (Hochachka and Somero 2002). Water temperature also has a significant influence on nerve, endocrine and reproductive systems (Luksiene and Svedang 1997). Interestingly, despite all of these potential impacts several medium and long-term monitoring studies (Holbrook et al. 1994, Malcolm et al. 2007; Stuart-Smith et al. 2010; James et al. 2012) have indicated that the composition of many coastal fish assemblages has remained relatively stable, despite increases in water temperature. One commonality amongst the abovementioned fish assemblages was that they were dominated by resident species.

Unlike migratory fish species, resident fishes have evolved to be tolerant of fluctuating diel and seasonal temperature regimes and many have therefore developed a broad physiological tolerance to temperature. Pörtner and Knust (2007), using laboratory studies on the eelpout (Zoarces viviparous), identified oxygen limitation as the primary mechanism driving thermally induced distributional shifts in fishes. Based on the assumption that this is the primary mechanism, scientists can predict the spatio-temporal distributional patterns of fishes using a combination of respirometry experiments (to determine the upper and lower thermal limits of species) and water temperature forecasting. However, with the evolution of eurythermy in many resident fishes, oxygen limitation may not necessary be a primary mechanism driving shifts, except at the margins of their distribution ranges. In these cases other mechanisms such as changes in reproductive output may drive distributional shifts. For example, reproductive failure in an area that becomes unfavourable, or the successful recruitment of eggs and larvae into a new favourable environment (Figueira et al. 2009) may drive distributional shifts in eurythermic resident species.

Water temperature has a major influence on many aspects of fish reproduction including fecundity and egg size (e.g. Brown et al. 2006), the rate of egg development (Hilder and Pankhurst 2003; Rombough 1997; Taranger and Hansen 1993), gonad maturation (Davies and Hanyu 1986), and the timing (Lobel 1989; Pankhurst and Porter 2003; Sheaves 2006) and frequency (Hilder and Pankhurst 2003; Pankhurst and Porter 2003) of reproductive activity. Besides water temperature, the timing of reproductive activity can also be influenced by a combination of other ecological drivers as well as physiological and phylogenetic factors, which can operate either through their effect on gametes, larvae, juveniles and/or adults (Sheaves 2006). The timing of reproduction may be geared to minimize egg predation, maximize dispersal, or provide larvae with maximum opportunity to survive in waters with patchy and irregular distribution of food (Shapiro et al. 1988). Reproductive timing in fishes is governed by proximate causal factors which serve to cue gonad development in adult fishes and ultimate causal factors which have developed through natural selection acting on survivorship and recruitment of larvae (Lobel 1989). Water temperature has long been recognised as an important proximate reproductive cue in fishes (Lobel 1989; Shapiro et al. 1988), but may also be an ultimate factor as it plays a role in determining the reproductive episodes (reviewed in Pankhurst and Porter 2003).

The reproduction of commercially important resident species such as those belonging to the families Sparidae, Lutjanidae and Serranidae has been reviewed by several authors (Robertson 1991; Sadovy 1996; Sheaves 2006). Many of the species belonging to these families have adopted a bet-hedging spawning strategy (asynchronous spawning) over an extended season to account for variability in egg and larval survival (Robertson 1991). Water temperature was identified as a major determinant of the spawning season in the Sparidae (Sheaves 2006). However, seasonality amongst the other families was thought to be determined by a complex of variables which are influenced by geography and adult biology (Robertson 1991; Sadovy 1996). Water temperature was identified as a potential factor influencing spawning periodicity and Sadovy (1996) concluded that the approximate spawning temperature range of resident reef fishes (excluding sparids) is generally narrower $(2-3^{\circ})$ in many tropical regions, when compared with more temperate habitats.

With clear relationships between temperature and spawning periodicity, it follows that ocean warming may alter the reproductive scope of resident fishes, which could in turn, result in distributional shifts. Reproductive scope in this manuscript is defined as the "breadth" of opportunity for reproduction and is determined as the area and time with suitable environmental and biological conditions including inter alia temperature, photoperiod, pH, nutrient availability and access to mates, for spawning. It is a multidimensional index that includes time, space and the importance of any specific spatial, temporal or environmental variable can be easy investigated by integrating the index over all other variables. For example, given a temperature range suitable for spawning by a particular species, the total area suitable for spawning can be estimated by simply integrating over months of the year. Alternatively by integrating over area, the time suitable for spawning during a calendar year can be determined. From a water temperature perspective, ocean warming hotspots provide ideal observatories for the investigation of the impacts of warming on the reproductive scope of fishes. These areas, characterised by above average ocean warming, have been identified in many areas including the northern Benguela (Monteiro et al. 2008) and eastern Australia (Byrne et al. 2009), and the response of organisms could provide predictive information for areas that are warming at normal rates.

The Diplodus sargus sub-species complex inhabits the shallow areas (<20 m depth) of the continental shelf in the warm-temperate waters of the Mediterranean, Northeast Atlantic, Southeast Atlantic, Southwest Atlantic, and Western Indian Ocean. The complex is economically valuable to the inshore recreational (Pradervand 2004; Mann and Buxton 1992), subsistence (Richardson et al. 2011a), artisanal (Pajuelo and Lorenzo 2002) and commercial fisheries (Morales-Nin et al. 2005). Like most sparids, this complex is reef associated and displays highly resident behaviour (Erzini and Abecasis 2009; D'Anna et al. 2011; Watt-Pringle 2009). Members of the complex have been described as rudimentary (Richardson et al. 2011b) and protandrous hermaphrodites (Mann and Buxton 1998) suggesting that they demonstrate some flexibility in their reproductive style. Despite this flexibility, all sub-species are asynchronous spawners, have protracted reproductive seasons (up to 5 months) during which they spawn regularly, even daily and consequently have a high annual egg production (Mylonas et al. 2011). After spawning in shallow waters (<6.0 m depth, Richardson 2010), the buoyant, pelagic eggs of species in this complex hatch after 50 to 90 h depending on ambient temperature (Divanach et al. 1982), and larvae remain pelagic for between 14–17 days (Di Franco et al. 2012). Larval dispersal is thought to be relatively wide (100–200 km scale) and post-settlement dispersal of at least 30 km was also observed (Di Franco et al. 2012). There is no information on the thermal tolerance of their eggs and larvae.

This study aims to assess the impact of ocean warming in the northern Benguela "hotspot" on the reproductive scope of an economically important coastal resident fish species *D. s. capensis*. Specifically, the objectives were to investigate the relationship between the water temperature in the well mixed shallow waters (<6.0 m depth) and reproductive activity in southern Angola, conduct a metadata analysis on the relationship between water temperature and reproductive activity within the *D. sargus* complex, and use these data to understand the recent trends in the distribution and duration of suitable thermal conditions throughout their northern Benguela distribution. The findings of the study are then discussed in the context of resident species worldwide.

Materials and methods

Sampling

Sampling was conducted between April 2008 and March 2009 in southern Angola between the towns of Namibe and Tômbua (Fig. 1). Between 29 and 61 adult fish were collected monthly by shallow water spearfishing (<6.0 m depth), hook-and-line techniques from the shore, or purchased from local markets. Each fish was measured (1 mm) (fork length), weighed (whole and without viscera) (0.1 g), and sexed. The gonads were weighed (0.1 g) and the state of gonad maturation was assessed macroscopically (Table 1). Sea surface temperature (SST) was measured daily in situ, whenever possible, for the duration of the study.

Data analysis

Reproductive seasonality was investigated by comparing both the proportion of female fish with ripe gonads (Table 1) in each monthly sample, and by examining peaks in the mean monthly female gonadosomatic index (GSI). A non-linear step model was used to examine the relationship between the

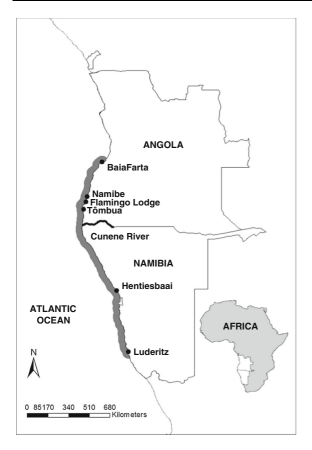


Fig. 1 Map showing the present known distribution of *Diplodus sargus capensis* on the west coast of southern Africa and the locations mentioned in the text

proportion of females that are ripe and SST. The stepmodel has the form:

$$PR = \begin{cases} a & \text{if } T > d \\ b + cT & \text{otherwise} \end{cases}$$

where *PR* is the proportion of ripe females, and *a*,*b* and *c* are estimated coefficients with *d* being a critical temperature. In the model $d = \frac{a-b}{c}$ is not estimated. Parameter estimates were obtained using non-linear least squares while parameter variability was estimated using parametric bootstrapping with a condition that the proportion of ripe females cannot be below zero.

Least-squares linear regression was used to examine the relationship between GSI and both photoperiod and mean monthly SST. Logistic regression was used to examine the statistical relationship between reproductively ripe fish and both photoperiod and mean monthly SST.
 Table 1
 Staging criteria used to macroscopically assess the gonads of *Diplodus sargus capensis* in southern Angola (validated microscopically in Richardson 2010)

Stage	Macroscopic description Ovaries are light with no oocytes visible to the naked eye. Testes increase in size and appear as flat white bands. No sperm present in tissue				
Resting					
Active	Ovaries increase in diameter and oocytes become clearly visible. Ovaries are orange/yellow. Veins and arteries become visible on ovaries. Testes increase in size taking on a triangular shape in cross-section, creamy-beige in colour and sperm present in the tissue when severed and pressure is applied				
Ripe	Ovaries large in diameter and yellow. Oocytes large and occupy the entire gonad and veins and arteries large and plentiful. Testes large and creamy white in colour, with sperm plentiful if the testes are cut and gently squeezed				
Spent	Ovaries bloodshot, flaccid and sac-like with few vitellogenic oocytes visible. Testes shrivelled, reduced in size and grey in colour				

Reproductive seasonality in the *Diplodus sargus* "complex"

Available published information on the spawning activity of fishes belonging to the *D. sargus* complex was collated. The months of observed spawning activity and the average monthly SST associated with those months was noted. In cases where average monthly SST was not available in the Mediterranean, the monthly SST climatology for that location was obtained from http://emis.jrc.ec.europa.eu/emis_3_1.php.

Trends in sea surface temperature

Long-term monthly SST data with a pixel resolution of 4 km^2 were obtained for the present known distribution of *D. s. capensis*, which extends from Lüderitz ($26^{\circ}38'47''S 15^{\circ}08'58''E$) in southern Namibia to Baia Farta ($12^{\circ}36'31''S 13^{\circ}12'04''E$) in northern Angola (Fig. 1), from 1982 to 2007 using the AVHRR Pathfinder 5.2 (Casey et al. 2010) dataset. All geore-ferenced pixels adjacent to the seaward side of the landmask were extracted for each month and each year.

Trends in reproductive scope

Total annual reproductive scope by area was defined as the total potential reproductive area per year and was calculated as the sum of all of the georeferenced SST pixels (each pixel represents 4 km²) that fell within the spawning temperature range of the subspecies complex throughout the Angolan and Namibian distribution of the species. The duration of the spawning season, or seasonal reproductive scope, was calculated as the number of months with SST pixels that were within the spawning temperature range and was calculated for localities with significant fisheries that targeted this species-Namibe, Angola (15°11'07"S 12°06'44"E) and Hentiesbaai, Namibia (22°07′16″S 14°16′27″E). Trends in reproductive scope (by area and season) between 1982 and 2008 were investigated using linear regression with year as the independent variable.

Results

Angola in situ water temperatures

An average of 22 SST measurements were recorded per month. The monthly means ranged from 16.5° to 24.7° C and the mean temperature was 20.4° C for the duration of the study period. There was a clear seasonal pattern with water temperature decreasing in May and remaining cold from June to October after which a sharp increase was observed in November (Fig. 2, top panel). An anomalous drop in the mean water temperature was observed during January.

Reproductive seasonality

There was a clear increase in the proportion of ripe mature females during the cold water months from June to October (Fig. 2, middle panel). Female reproductive activity was low during summer, however, a small increase in the proportion of ripe individuals was observed during January and February. This pattern was mirrored by the GSI information (Fig. 2, bottom panel). *D. s. capensis* with ripe running gonads were observed during months with mean SST of between 16 and 20 °C.

The step-model estimated that the proportion of ripe females declined rapidly until 19.33 °C (95 % CI = 19.0–20.0 °C) after which the proportion stabilised at 0.08 (Fig. 3, top panel). The step-model explained 94 % of the variation within the data.. Significant relationships were noted between female

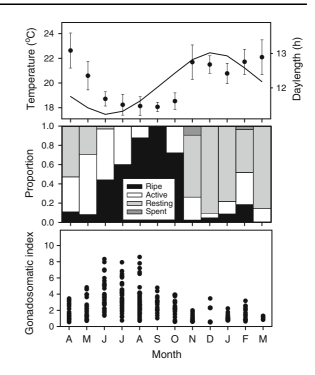


Fig. 2 Mean (\pm SD) surf zone water temperature (n = 269) and daylength (*top panel*), percentage of mature fish in different macroscopic stages (n = 578) (*middle panel*), and individual gonadosomatic indices (n = 571) (*bottom panel*) for mature female *Diplodus sargus capensis* from southern Angola sampled between April 2008 and March 2009

GSI and both photoperiod (T = -7.12, P < 0.01) and mean SST (T = 8.89, P < 0.01) (Fig. 3, bottom panel). In the case of the monthly proportion of ripe females, only mean SST was found to be statistically significant (P < 0.01) (Fig. 3, middle panel). The variance of the data explained by the regression models ranged between $R^2 = 0.26$ and 0.39, respectively.

Metadata analysis

The reproductive biology of the *D. sargus* complex has been studied in the Mediterranean, North Atlantic, South Atlantic, Western Indian Ocean and Persian Gulf (Table 2). A total of 13 studies (including this one) provided information on the spawning periodicity of 14 populations. These populations occurred in areas with monthly average temperatures ranging from 14 to 26 °C. Spawning of *D. sargus* complex occurred between 15 and 20 °C regardless of their location and ambient temperature regime (Table 2).

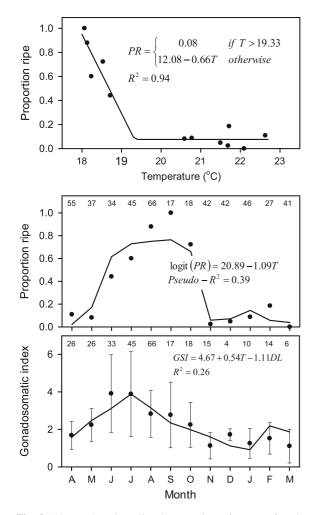


Fig. 3 Observed and predicted proportion of mature female fish with ripe gonads (*top panel*) as a function of mean surf zone water temperature, and the observed and predicted proportion of mature female fish (*middle panel*) and mean (\pm SD) gonadosomatic index (*bottom panel*) for *Diplodus sargus capensis* sampled from southern Angola between April 2008 and March 2009. Sample sizes are included at the top of each panel. *PR* proportion ripe, *GSI* gonadosomatix index, *T* mean surfzone temperature, *DL* day length

Changes in reproductive scope

There was been a general increase in the water temperatures of the northern Benguela region over the last three decades and based on spawning temperatures between 15 and 20 °C, there was a significant (P < 0.01) 134 km² (0.7 %) reduction in the annual reproductive scope by area in Angola (Fig. 4, top panel). In contrast, there was a significant (P < 0.01) increase of 249 km² (0.6 %), despite possible cyclical trends, in the annual reproductive scope by area in

Namibia (Fig. 4, bottom panel). The reproductive scope of *D. s. capensis* by season, decreased on average by 1.05 months per decade (P < 0.01) in the waters of Namibe, Angola compared with an insignificant (P = 0.08) increase of 0.76 months per decade in Hentiesbaai, Namibia (Fig. 5). The decreased reproductive scope by season in southern Angola and an increasing trend in northern Namibia is illustrated in Fig. 6 with the Cunene River mouth region (approximately -17° S), the transition zone.

Discussion

Peak reproductive activity for D. sargus capensis in Angola coincided with the lowest recorded water temperatures during the study period (June to October) and was found to be significantly correlated with water temperature. The results of the metadata analysis (Table 2) supported these findings, but also showed that regardless of the ambient temperatures, populations in this species complex spawn when the temperatures range between 15° and 20 °C. It appeared that the season (or photoperiod) did not have a major influence on the timing of reproduction in this species. The reasons for the rigid reproductive thermal range are unclear. Being resident, adult D. sargus capensis are obviously eurythermic (Table 2) and thus one may expect adults to have a broad reproductive thermal range. However, Sheaves (2006) found that sparids with a tropical distribution reproduce during the coolest period of the year and concluded that it was because of the cool water stenothermy of the early life stages. Generally, gametes, eggs and larvae are more sensitive to environmental fluctuations than adult fishes and have a narrower thermal tolerance (Pörtner and Farrell 2008; Pankhurst and Munday 2011). While there is no specific information on the relationship between larval survival and temperature in the Diplodus, Houde's (1989) review of the effects of water temperature on fish larval growth and mortality showed that mortality increased by 0.01 with a 1 °C increase in water temperature. The optimal temperature range for adult spawning activity amongst the D. sargus sub-species complex, probably has, like in other fishes (Lobel 1989), been influenced by natural selection through the survival of gametes, eggs and larvae.

The reproductive stenothermy makes the *D. sargus* sub-species complex extremely susceptible to ocean

Species	Location, ocean	Spawning period	Spawning duration (months)	Spawning temperature (°C)	Temperature at location (°C)
D. sargus cadenati ¹	Canary Islands, Northeast Atlantic	Dec-Apr	5	18–20	18–25
D. sargus cadenati ²	Mauritania, Northeast Atlantic	Jan–Apr	4	16-20*	16–26*
D. sargus capensis ³	Eastern Cape, Southwest Indian	May-Dec	8	16-20*	16-22*
D. sargus capensis ⁴	KwaZulu-Natal, Southwest Indian	May-Dec	8	17-20*	17–24*
D. sargus capensis ⁵	Western Cape, Southwest Indian	Aug–Mar	5	17-20*	16-22*
D. sargus capensis ⁶	Southern Angola, Southeast Atlantic	Jun-Oct	5	16-20	16–25
D. sargus sargus ⁷	Algeria, Western Mediterranean	Feb–Apr	3	15-18*	15-24*
D. sargus sargus ⁸	Azores, North Atlantic	Mar–Jun	4	15–17	14–24
D. sargus sargus ⁸	Lisbon, Northeast Atlantic	Feb-May	4	16–17*	15-20*
D. sargus sargus ⁹	Bay of Biscay, Northeast Atlantic	Apr–Jun	3	13–17	12–21
D. sargus kotschyi ¹⁰	Kuwait, Persian Gulf	Nov–Mar	5		
D. sargus sargus ¹¹	Egypt, Eastern Mediterranean	Jan–Apr	4	17-20*	17–27*
D. sargus sargus ¹²	Gulf of Lion, Western Mediterranean	Apr–May	2	14–16*	14-20*
D. sargus sargus ¹³	Gulf of Tunis, Western Mediterranean	Mar–May	3	15-18	14–25

Table 2 Comparison of the spawning period, duration, temperature and ambient temperatures of *Diplodus sargus* complex throughout the world

*Indicates an estimation based on the sea surface temperature climatology obtained from http://emis.jrc.ec.europa.eu/emis_3_1.php 1 = Pajuelo and Lorenzo (2004), 2 = Dia et al. (2009), 3 = Lasiak (1981), 4 = Joubert (1981), 5 = Mann and Buxton (1998), 6 = this study, 7 = Bencgalel and Kara (2010), 8 = Morato et al. (2003), 9 = Lavín and Cabanas (2000), 10 = Abou-Seedo et al. (1990), 11 = El Maghraby et al. (1982), 12 = Man-Wai (1985), 13 = Mouine et al. (2007)

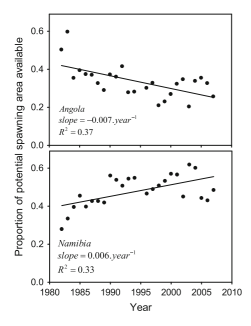


Fig. 4 Total annual reproductive scope of *Diplodus sargus capensis* from Angola and Namibia between 1982 and 2009. Data presented is the area available to spawn per year with temperatures between 15 and 20 °C expressed as a proportion of all area in the study area

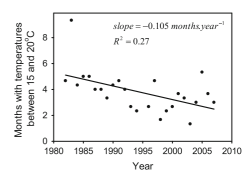


Fig. 5 Seasonal reproductive scope in terms of the duration of the spawning season (months with temperatures between 15 and 20 °C) for *Diplodus sargus capensis* from Namibe, southern Angola between 1982 and 2009

warming, but also allowed the correlative analysis of the trends in reproductive scope. This study showed a decline in the reproductive scope by area of approximately 7 %/decade in Angola. More startling perhaps was the one month per decade reduction in reproductive scope by season in Angola during the study period. As an asynchronous spawning fish that can spawn daily (Mylonas et al. 2011) during favourable

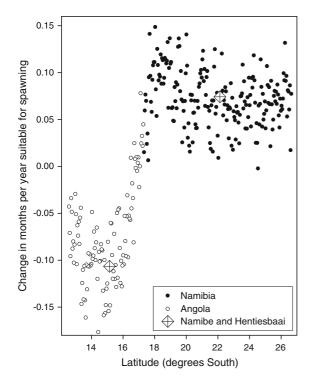


Fig. 6 Annual change in reproductive scope in terms of the duration of the spawning season (months with temperatures between 15 and 20 °C) of *Diplodus sargus capensis* from southern Angola to northern Namibia. Each point represents the slope from a linear regression of months suitable from spawning each year for each pixel (4 km²) against year between 1982 and 2009. The positions of Namibe in southern Angola and Hentiesbaai in northern Namibia are shown and correspond to the results illustrated in Fig. 5

conditions (in this case temperatures between 15 and 20 C), this equates to a reduction in reproductive output of approximately 20 % per decade.

If this trend was to continue, the reproduction of *D*. *s. capensis* will cease in Angola within 60 years. Although this may ultimately result in its extirpation in Angola and what we would perceive to be a "distributional shift" of the species, two possible scenarios may prevent this from happening. Firstly, larval connectivity from the Namibian breeding population, which showed a concomitant increase in reproductive scope, may result in the recruitment of juveniles into southern Angola. Although the hydrography of the Mediterranean is not comparable, Di Franco et al. (2012) showed that the pelagic larval dispersal of *D*. *s. sargus* extended at a scale of between 100 and 200 km. This suggests that at least some recruitment into Angola is possible. However, pelagic larval dispersal

into Angola would require favourable current transport and depend on the availability of suitable temperatures for larval survival. There is very little information on the speed and direction of the coastal currents in northern Namibia. However, Mouton et al. (2001) observed a consistent northward coastal current flow of over 12 cm s⁻¹ during July (which is within the reproductive scope of *D. s. capensis*) in northern Namibia. Although this suggests that the transport mechanism may be available for the eggs and larvae, there is no information to suggest that *D. sargus capensis* eggs and larvae would survive in Angolan waters, when the average water temperatures would exceed 20 °C, if the warming trend continues.

The second scenario that may prevent a local extinction in Angola and a "distribution shift" would be episodic reproductive events during periods with unusually cold water temperatures. As *D. sargus capensis*, like most resident fishes is long-lived, cold temperature anomalies (such as in January during this study) may provide opportunities for reproduction during their lifetime. The maximum age of *D. sargus capensis* in Angola was 20 years and the age at maturity was just under 2 years (Richardson et al. 2011a). Therefore, it is quite conceivable that temperatures favourable for spawning would arise during that 18 year reproductive lifespan of an individual and the population would persist.

The results indicated that the effect of ocean warming was not uniform through the distribution of the species. In contrast to Angola, Namibia was shown to have a 6 % per decade increase in reproductive scope by area, but there appeared to be fluctuations associated with the variable oceanographic conditions in Namibia (Hutchings et al. 2009) within each decade. The primarily southward extension of the reproductive scope by area suggested that reproduction is now possible further south. Therefore, if egg and larval dispersal patterns allow juvenile recruitment, these areas may provide conditions for the southward expansion of the *D. s. capensis* population, ultimately resulting in a distributional shift.

Although changes in reproductive scope may drive distribution shifts in resident species, under the present rates of ocean warming, this mechanism is unlikely to drive the rapid, large-scale distributional shifts predicted for migratory species, where oxygen limitation is the driving mechanism. However, the poleward nature of the shifts predicted for migratory fishes is also the most likely scenario for resident fishes. Perhaps more importantly, this study has shown that ocean warming may influence the seasonal reproductive scope of resident species and may be a primary mechanism driving variations in the abundance of fishes within their known distribution ranges. Based on these results, it appears that these variations will not be homogenous and ultimately some fisheries may benefit while others will experience declines in fish abundance. For example, in the city of Namibe (Fig. 1) and its close surrounds (4 km north and south of the town), D. s. capensis is overexploited (SBR = 20 %, Richardson 2010) and accounts for 82 % of the catch in the subsistence fishery (WMP, unpublished data). With the present one month per decade reduction in the seasonal reproductive scope (Fig. 5, 6), this overexploited population is unlikely to recover even with a reduction in fishing pressure. Therefore, ocean warming is likely to lead to social hardship in this community.

Although the relationship between reproductive output and recruitment success is far from clear (Dixon et al. 1999), an increase in the seasonal reproductive scope of this species in Namibia is expected to lead to improved recruitment success as egg production is considered to be a better indicator of recruitment potential than spawner biomass (Marshall et al. 1998; Sadovy 1996). While there is no significant subsistence fishery for this species, it was the fourth most important teleost species in Namibia (Kirchner et al. 2000) and an increase in recruitment success may ultimately provide improved recreational fishing opportunities and as an important tourist activity, indirect economic gains for the coastal communities of this region.

Sadovy (1996) found that the spawning of reef fishes (excluding sparids) occurred at temperatures below the maximum annual temperatures in many tropical and subtropical regions, such as Kuwait and the Red Sea, Hawaii, the Caribbean, Belau New Caledonia, Fiji and northern Australia. Donelson et al. (2010) found that the reproductive activity of a tropical damselfish (*Acanthochromis polyacanthus*) decreased or ceased in an elevated water temperature regime. Based on these studies, it is likely that other resident fishes would respond in a similar way to the *D. s. capensis* population and one could therefore expect a reduction in the reproductive scope (by area and season) of resident fishes in their tropical distributions, and a concomitant increase in the reproductive scope of these species in their more temperate distributions. However, in extreme cases, where resident fishes have extremely narrow reproductive temperature range, such as the Nassau grouper (*Epinephelus striatus*) which only spawn at temperatures between 25° and 26 °C (Tucker et al. 1993), the reproductive scope may decrease rapidly, and unless larval distribution allows juveniles of these species to settle in areas with suitable spawning temperatures, extinctions are possible.

One important consideration when predicting the impact of ocean warming on reproductive scope of resident fishes is the adaptability of some species to different thermal regimes (Pankhurst and Munday 2011). Local adaptations to thermal regimes have been observed in the populations of several species (Farrell et al. 2008; Munday et al. 2008). Thermal adaptability is generally considered to be dependent on the genetic variability associated with this characteristic and species with substantial genetic variability should be more adaptable to the affects of rapid warming. To predict the response to ocean warming the adaptability of species of interest should first be evaluated. Since a candidate gene approach would be lengthy and prohibitive, in the interim, a metadata analysis (as conducted in this study), that documents the reproductive thermal range amongst geographically distinct populations is probably the most appropriate method. However, in the absence of published information, laboratory experiments (see Donelson et al. 2010) or a metadata analysis of a closely related species may provide the necessary additional information.

The methods used in this study can be extended to evaluate some of the threats and opportunities facing fisheries that target resident fish species globally. Due to the heterogeneity of the expected changes demonstrated in this study, fisheries managers are likely to face significant challenges, and the requirement to evaluate the impact of changing ocean temperatures on the reproduction of resident fishes in their area. Our results suggest that this may be carried out using a broad range of methods. However, in the absence of an evaluation, some general trends can be expected. A reduction in the abundance or even extirpations can be expected on the equatorial range boundaries of resident fishes. This could be accompanied by a poleward expansion of the populations. On the whole, tropical resident species are expected, to shift into warm-temperate regions. This could, to some degree, mitigate for the loss of the warm-temperate species, which may move poleward into the cool-temperate zone. Unfortunately, the reductions in the abundance of resident fish in the tropics are unlikely to be mitigated by the "arrival" of any other species and fishing communities in these areas may face hardships in the future. These heterogeneous changes in fish abundance may be accompanied by unexpected changes in the dynamics of fisheries, such as the transboundary shift in the distribution of fishing effort, which may occur due to a perceived "distributional shift" of the target species. Regional Fisheries Management Organisations will have a significant management role in these circumstances and additional fishing effort in certain areas will have to be well monitored and controlled using appropriate catch and/ or effort controls.

Acknowledgments This material is based upon work supported by the National Research Foundation (NRF) under grant number 66957. Rico Sakko is thanked for the use of the facilities at Flamingo Lodge, Angola. Barry Wareham (Basil Manning Active Fishing Gear) is thanked for the fishing tackle sponsorship. TJR received financial support from Rhodes University, the Ernst and Ethel Eriksen Trust and the NRF. Amber Childs is thanked for her assistance with the map. These data were provided by the Group for High Resolution Sea Surface Temperature (GHRSST) and the US National Oceanographic Data Center. This project was supported in part by a grant from the NOAA Climate Data Record (CDR) Program for satellites. Two anonymous reviewers are thanked for their comments on an earlier version of this manuscript.

References

- Abou-Seedo F, Wright JM, Clayton DA (1990) Aspects of the biology of Diplodus sargus kotschyi (Sparidae) from Kuwait Bay. Cybium 14:217-223
- Bencgalel W, Kara MH (2010) Age, croissance et reproduction du sar commun Diplodus sargus sargus (Sparidae) des Cotes de l'est Algerien. Rapp Comm int Mer Medit 39:451
- Brown NP, Shields RJ, Bromage NR (2006) The influence of water temperature on spawning patterns and egg quality in the Atlantic halibut (Hippoglossus hippoglossus L.). Aquaculture 261:993-1002
- Byrne M, Ho M, Selvakumaraswamy P, Nguyen HD, Dworjanyn SA, Davis AR (2009) Temperature, but not pH, compromises sea urchin fertilization and early development under near-future climate change scenarios. Proc R Soc 276:1883-1888
- Casey KS, Brandon TB, Cornillon P, Evans R (2010) The past, present and future of the AVHRR pathfinder SST program. In: Gower JFR, Alberotanza L, Barale V (eds) Oceanography from space. Springer, New York

D'Anna G, Giacalone VM, Pipitone C, Badalamenti F (2011) Movement pattern of white seabream, Diplodus sargus (L., 1758) (Osteichthyes, Sparidae) acoustically tracked in an artificial reef area. Ital J Zool 78:255-263

Rev Fish Biol Fisheries (2014) 24:493-504

- Davies PR, Hanyu I (1986) Effect of temperature and photoperiod on sexual maturation and spawning of the common carp: I. Under conditions of high temperature. Aquaculture 51:277-288
- Di Franco A, Gillanders BM, De Benedetto G, Pennetta A, De Leo GA, Guidetti P (2012) Dispersal patterns of coastal fish: implications for designing networks of marine protected areas. PLoS ONE 7(2):e31681. doi:10.1371/journal. pone.0031681
- Dia M, Wague A, Ghorbel M (2009) Etude de la reproduction chez Diplodus sargus cadenati (Sparidae) de la Cote de Nouadhibou (Mauritanie). Bull Inst Natn Scien Tech Mer de Salammbô 36:21-30
- Divanach P, Kentouri M, Paris J (1982) Etapes du developpement embryonnaire et larvaire du sar, Diplodus sargus L., en elevage. Aquaculture 27:339-353
- Dixon PA, Milicich MJ, Sugihara G (1999) Episodic fluctuations in larval supply. Science 283:1528-1530
- Donelson JM, Munday PL, McCormick MI, Pankhurrt NW, Pankhurst PM (2010) Effects of elevated water temperature and food availability on the reproductive performance of a coral reef fish. Mar Ecol Prog Ser 401:233-243
- El Maghraby AM, Botros GA, Hashem MT, Wassef EA (1982) Maturation, spawning and fecundity of two sparid fish Diplodus sargus, L. and Diplodus vulgaris, Geoff. in the Egyptian Mediterranean waters. Bulletin Bull Inst Oceanogr Fish ARE 8:51-67
- Erzini K, Abecasis D (2009) Home range, residency and movements of Diplodus sargus and Diplodus vulgaris in a coastal lagoon: connectivity between nursery and adult habitats. Estuar Coast Shelf Sci 85:525-529
- Farrell AP, Hinch SG, Cooke SJ, Patterson DA, Crossin GT, Lapointe M, Mathes MT (2008) Pacific salmon in hot water: applying aerobic scope models and biotelemetry to predict the success of spawning migrations. Physiol Biochem Zool 81:697-708
- Figueira WF, Biro P, Booth DJ, Valenzuela VC (2009) Performance of tropical fishes recruiting into temperate habitats: role of ambient temperature and implications of climate change. Mar Ecol Prog Ser 384:231-239
- Hilder ML, Pankhurst NW (2003) Evidence that temperature change cues reproductive development in the spiny damselfish, Acanthochromis polyacanthus. Environ Biol Fishes 66:187-196
- Hochachka PW, Somero GN (2002) Biochemical adaptation: mechanism and process in physiological evolution. Oxford University Press, New York
- Holbrook SJ, Kingsford MJ, Schmitt RJ, Stephens JS (1994) Spatial and temporal patterns in assemblages of temperate reef fish. Am Zool 34:463-475
- Houde E (1989) Comparative growth, mortality and energetics of marine fish larvae: temperature and implied latitudinal effect. Fish Bull US 87:4712-4950
- Hutchings L, van der Lingen CD, Shannon LJ, Crawford RJM, Verheye HMS, Bartholomae CH, van der Plas AK, Louw D, Kreiner A, Ostrowski M, Fidel Q, Barlow RG, Lamont T, Coetzee J, Shillington F, Veitch J, Currie JC, Monteiro

PMS (2009) The Benguela current: an ecosystem of four components. Prog Oceanogr 83:15–32

- James NC, Götz A, Cowley PD, Potts WM (2012) Temporal variability of a temperate fish assemblage in Africa's oldest marine protected area. Afr J Mar Sci 34:15–26
- Joubert CSW (1981) Aspects of the biology of five species of inshore reef fishes on the Natal coast, South Africa. Invest Rep Oceanogr Res Inst 51:1–16
- Kirchner CH, Sakko AL, Barnes JI (2000) An economic valuation of the Namibian shore-angling fishery. S Afr J Mar Sci 22:17–25
- Lasiak TA (1981) Nursery grounds of juvenile teleosts: evidence from the surf zone of Kings Beach, Port Elizabeth. S Afr J Sci 77:388–390
- Last PR, White WT, Gledhill DC, Hobday AJ, Brown R, Edgar GJ, Pecl G (2011) Long-term shifts in abundance and distribution of a temperate fish fauna: a response to climate change and fishing practices. Glob Ecol Biogeogr 20:58–72
- Lavín A, Cabanas JM (2000) Spanish standard sections. Area 4 (Bay of Biscay and Eastern Atlantic) Spanish Report. In: Turrel B (ed) The annual ICES ocean climate status summary 1999/2000 ICES environmental status reports. Copenhagen: International Council for the Exploration of the Sea, pp 1–8 (Annex J.)
- Lobel PS (1989) Ocean current variability and the spawning season of Hawaiian reef fishes. Environ Biol Fishes 24: 161–171
- Luksiene D, Svedang H (1997) A review on fish reproduction with special reference to temperature anomalies. Fiskeriverket, Fiskeriverket, Kustlaboratoriet, Ö regrund
- Malcolm HA, Gladstone W, Lindfield S, Wraith J, Lynch TP (2007) Spatial and temporal variation in reef fish assemblages of marine parks in New South Wales, Australia baited video observations. Mar Ecol Prog Ser 350:277–290
- Mann BQ, Buxton CD (1992) Diets of *Diplodus sargus capensis* and *Diplodus cervinus hottentotus* (Pisces: Sparidae) on the Tsitsikamma coast, South Africa. Koedoe 35:27–36
- Mann BQ, Buxton CD (1998) The reproductive biology of Diplodus sargus capensis and D. cervinus hottentotus (Sparidae) off the south-east Cape coast, South Africa. Cybium 22:31–47
- Man-Wai R (1985) Les sars du golfe du Lion, Diplodus sargus, D.vulgaris, D. annularis (Pisces, Sparidae). Ecobiologie, Pêche. Dissertation, University of Montpellier
- Marshall CT, Kjesbu OS, Yaragina NA, Solemday P, Ulltang Ø (1998) Is spawner biomass a sensitive measure of the reproductive and recruitment potential of Northeast Arctic cod? Can J Fish Aquat Sci 55:1766–1783
- McFarlane GA, King JR, Beamish RJ (2000) Have there been recent changes in climate? Ask the fish. Prog Oceanogr 47:147–169
- Monteiro PMS, van der Plas AK, Me'liced JL, Florenchie P (2008) Interannual hypoxia variability in a coastal upwelling system: ocean–shelf exchange, climate and ecosystem-state implications. Deep-Sea Res 55:435–450
- Morales-Nin B, Moranta J, García C, Tugores MP, Grau AM, Riera F, Cerdà M (2005) The recreational fishery off Majorca Island (western Mediterranean): some implications for coastal resource management. ICES J Mar Sci 62:727–739

- Morato T, Afonso P, Lourinho P, Nash RDM, Santos RS (2003) Reproductive biology and recruitment of the white seabream in the Azores. J Fish Biol 63:59–72
- Mouine N, Francour P, Ktari M, Chakroun-Marzouk N (2007) The reproductive biology of *Diplodus sargus sargus* in the Gulf of Tunis (central Mediterranean). Scientia Marina 71:461–469
- Mouton DP, Boyd AJ, Bartholomae CH (2001) Near-surface currents and hydrology off northern Namibia in July 1999. S Afr J Sci 97:209–212
- Munday PL, Jones GP, Pratchett MS, Williams AJ (2008) Climate change and the future for coral reef fishes. Fish Fish 9:261–285
- Mylonas CC, Zohar Y, Pankhurst N, Kagawa H (2011) Reproduction and broodstock management. In: Pavlidis M, Mylonas C (eds) Sparidae: biology and aquaculture of gilthead sea bream and other species. Wiley-Blackwell, Oxford
- Pajuelo JG, Lorenzo JM (2002) Growth and age estimation of Diplodus sargus cadenati (Sparidae) off the Canary Islands. Fish Res 59:93–100
- Pajuelo JG, Lorenzo JM (2004) Basic characteristics of the population dynamic and state of exploitation of Moroccan white seabream *Diplodus sargus cadenati* (Sparidae) in the Canarian archipelago. J Appl Ichthyol 20:15–21
- Pankhurst NW, Munday PL (2011) Effects of climate change on fish reproduction and early life history stages. Mar Freshw Res 62:1015–1026
- Pankhurst NW, Porter MJR (2003) Cold and dark or warm and light: variations on the theme of environmental control of reproduction. Fish Physiol Biochem 28:385–389
- Perry AL, Low PJ, Ellis JR, Reynolds JD (2005) Climate change and distribution shifts in marine fishes. Science 308: 1912–1915
- Pörtner HO, Farrell AP (2008) Physiology and climate change. Ecology 322:690–692
- Pörtner HO, Knust R (2007) Climate change affects marine fishes through the oxygen limitation of thermal tolerance. Science 315:95–97
- Pörtner HO, Peck MA (2010) Climate change effects on fishes and fisheries: towards a cause-and-effect understanding. J Fish Biol 77:1745–1779
- Pradervand R (2004) Long-term trends in the shore fishery of the Transkei coast, South Africa. Afr Zool 39:247–261
- Richardson TJ (2010) The taxonomy, life-history and population dynamics of blacktail, *Diplodus capensis* (Perciformes: Sparidae), in southern Angola. Dissertation, Rhodes University
- Richardson TJ, Potts WM, Santos CV, Sauer WH (2011a) Comparison of the population structure and life-history parameters of *Diplodus capensis* (Sparidae) in exploited and unexploited areas of southern Angola. Afr J Mar Sci 33:191–201
- Richardson TJ, Potts WM, Sauer WH (2011b) The reproductive style of *Diplodus capensis* (Sparidae) in southern Angola: rudimentary hermaphroditism or partial protandry? Afr J Mar Sci 33:321–326
- Robertson DR (1991) The role of adult biology in the timing of spawning of tropical reef fishes. In: Sale PF (ed) the ecology of fishes on coral reefs. Academic Press, San Diego, pp 356–386

- Rombough PJ (1997) The effects of temperature on embryonic and larval development. In: Wood CM, McDonald DG (eds) Global warming: implications for freshwater and marine fish. Cambridge University Press, Cambridge, pp 177–223
- Sadovy YJ (1996) Reproduction of reef fishery species. In: Polunin VC, Roberts CM (eds) Reef Fisheries. Chapman and Hall, London, pp 15–59
- Shapiro DY, Hensley DA, Appeldoorn RS (1988) Pelagic spawning and egg transport in coral-reef fishes: a skeptical overview. Environ Biol Fishes 22:3–14
- Sheaves M (2006) Is the timing of spawning in sparid fishes a response to sea temperature regimes? Coral Reefs 25: 655–669
- Stebbing ARD, Turk SMT, Wheeler A, Clarke KR (2002) Immigration of southern fish species to south-west England

linked to warming of the North Atlantic (1960–2001). J Mar Biol Assoc UK 82:177–180

- Stuart-Smith RD, Barrett NS, Stevenson DG, Edgar GJ (2010) Stability in temperate reef communities over a decadal time scale despite concurrent ocean warming. Glob Change Biol 16:122–134
- Taranger GL, Hansen T (1993) Ovulation and egg survival following exposure of Atlantic salmon, Salmo salar L., broodstock to different water temperatures. Aquac Res 24: 151–156
- Tucker JW, Bush PG, Slaybaugh ST (1993) Reproductive patterns of Caymen Islands Nassau grouper (*Epinephelus striatus*) populations. Bull Mar Sci 52:961–969
- Watt-Pringle PA (2009) Movement behaviour of three South African inshore sparid species in rocky intertidal and shallow subtidal habitats. Dissertation, Rhodes University