Latitudinal trends in habitat quality of shallow-water flatfish nurseries

Vânia Freitas¹,²,*, Sebastiaan A. L. M. Kooijman³, Henk W. van der Veer¹

¹Royal Netherlands Institute for Sea Research, PO Box 50, 1790 AB Den Burg, Texel, The Netherlands
²Centro Interdisciplinar de Investigação Marinha e Ambiental (CIIMAR), Universidade do Porto, Rua dos Bragas 289, 4050-123 Porto, Portugal
³Department of Theoretical Biology, Vrije Universiteit, de Boelelaan 1087, 1081 HV Amsterdam, The Netherlands

ABSTRACT: The habitat quality of European shallow-water nurseries was studied for 3 common flatfish species based on juvenile growth conditions. Field growth of 0-group plaice Pleuronectes platessa, flounder Platichthys flesus and sole Solea solea, from both published and unpublished studies, was compared with maximum growth predicted by a bioenergetics model based on the dynamic energy budget theory. In plaice and flounder, realized growth ratio decreased consistently during the growing season in most of the nurseries analyzed, indicating a widespread pattern of declining conditions. In sole, growth performance was not maximal, but as opposed to the other species, no clear temporal trend in realized growth ratio was observed. A latitudinal comparison of realized growth ratio over the various nurseries indicated clear positive trends for plaice and flounder, with better growth conditions at northern latitudes. In sole, despite some variability, the same trend was found during part of the summer. In the absence of clear gradients in benthic prey biomass, we hypothesize that increased food limitation in southern locations is most likely caused by interspecific competition reducing maximum individual intake rates. These results suggest that, in the context of global warming, habitat quality of southern European nurseries for juvenile growth may be particularly affected by the combined interaction of food and thermal constraints.

KEY WORDS: Plaice · Flounder · Sole · Growth potential · Dynamic energy budget · Temperature · Food limitation · Latitude

INTRODUCTION

Temperate estuarine and shallow coastal systems are key habitats in the context of life cycle closure in marine populations (sensu Sinclair 1988). For flatfish, these areas act as nurseries where the trade-off between growth and survival is optimized due to abundant food resources, few predators and favourable environmental conditions (Bergman et al. 1988b, Gibson et al. 2002). These specific habitat quality features in combination with the amount of area available have the potential to influence recruitment levels (Rijnsdorp et al. 1992, Gibson 1994, Van der Veer et al. 1994, 2000) and thus, over the years, there has been an increasing interest in better evaluating the function of nurseries (Vasconcelos et al. 2011). Although the production of juveniles to the adult populations has been suggested as the most integrative measure of the quality of juvenile habitats (Able et al. 1999, Beck et al. 2001), owing to problems in accurately estimating population sizes and their changes (Cowley & Whitfield 2002), other indicators have been sought. Growth is inextricably linked to mortality in the early life stage, which is supported by the general inverse relationship between size of an individual fish and its risk of mortality (Zijlstra et al. 1982, Beverton & Iles 1992, Sogard 1997). This has motivated the use of fish growth as a suitable index...

Classical studies with flatfish have explored the ability of fish to achieve its maximum growth potential as a relative measure of its well-being and, indirectly, of the quality of its coastal nurseries (Zijlstra et al. 1982, Van der Veer 1986, Van der Veer et al. 1991, 2001a). While studies from the western Dutch Wadden Sea indicated that growth in the field was generally maximal during most of the nursery phase, research in other areas showed a slightly different picture, with increasing discrepancies between observed field growth during summer and maximum growth measured in the laboratory (Amara 2004, Hurst & Abookire 2006, Freitas et al. 2010a). In plaice Pleuronectes platessa, such a trend was observed over a large latitudinal range from northern Norway estuarine waters (Freitas et al. 2010a), to the Irish Sea (Nash et al. 1994), and at Balgzand, in the Dutch Wadden Sea (Teal et al. 2008). Although this discrepancy suggested food limitation, an experimental bias could not be excluded since, in summer, fish size had outgrown the range where the maximum growth model could be applied (Van der Veer et al. 2010).

This impasse was recently solved with the use of a mechanistic growth model based on the dynamic energy budget (DEB) theory (Kooijman 2000, 2010). DEB models provide a benchmark for maximum growth potential that simultaneously considers both the internal state of the individual (structure, reserve) and external factors (e.g. food availability and temperature) affecting metabolism (Kooijman 2000). This modelling approach was applied to re-investigate 0-group plaice growth conditions in the Balgzand intertidal area (Van der Veer et al. 2010) and previous conclusions have been refined. Evidence from several years suggested a reduced habitat quality for plaice in summer, proposing competition with (or interference by) gobiodae and epibenthic crustaceans as the most likely underlying mechanism (Van der Veer et al. 2010).

At present, it is unclear to what extent these results for 0-group plaice at Balgzand are a general phenomenon present in more areas and whether other flatfish species exhibit similar patterns during their nursery phase. On the one hand, the same dominant mobile epibenthic fauna are present across temperate shallow-water systems (Pihl 1985, Maes et al. 1998, Amara & Paul 2003, Freitas et al. 2010a) with annual density maxima in summer (Maes et al. 1998, Henderson et al. 2006, Viegas et al. 2007), which could indicate that similar processes are likely to take place regardless of the area considered. On the other hand, over a large spatial scale, different patterns could emerge due to variations in temporal utilization of the nursery by juvenile flatfish (e.g. timing of nursery colonization). At the interspecific level, differences in feeding and environmental preferences may also offset potential competitive effects.

We evaluated these hypotheses by comparing the conditions for growth in different flatfish species and various nurseries along their range of distribution. This study builds on the approach followed in Van der Veer et al. (2010) to study plaice growth conditions using bioenergetics modelling and expands the scope to 2 other flatfish species, sole Solea solea and flounder Platichthys flesus. These two species are also important components of demersal flatfish communities, overlapping with plaice in the use of coastal nursery grounds from Bay of Biscay to Western Scotland (Nielsen 1986, Quéro et al. 1986). As far north as the White Sea (Nielsen 1986), only plaice and flounder can be found, while in southern Europe, sole and flounder co-occur up to a latitude of ~40°N (Vinagre et al. 2005).

MATERIALS AND METHODS

Field data

Basic data were collected from published and unpublished field studies describing juvenile flatfish growth in combination with local temperature conditions. Studies were selected to cover most of the species distribution ranges and those providing information for various species were particularly targeted. Information presented in graphical form (seasonal changes in mean total length and in water temperature) was scanned and digitalized using Grab-it! XP 10 software (Datatrend Software). Details on the sampling procedures of published data sets can be found in the papers (see Table 1 for references). In most areas sampling was performed with beam trawls (1 to 4 m), but in very shallow waters push nets and drop traps were also used.

Temperature data were as much as possible taken from the original publications; however, in some cases other sources were used to complement the data. These included published studies reporting seasonal temperature patterns for the same areas and years of study or oceanographic databases available from national research or meteorological institutes (see Table 1 for sources). The temperature conditions in each location considered in this study are shown in Appendix 1 (Fig. A1).
Field growth rates were estimated based on reported increases in mean total length (cm d⁻¹) of the entire population or specific age classes (or cohorts) between 2 successive sampling dates. To avoid the confounding effects of migrations on growth estimates, in each data set only the period of linear increase in mean length was considered assuming that: (1) during settlement (spring–summer), length–frequency distributions are biased by the continuous arrival of small individuals, resulting in an underestimation of growth rates; and (2) at the end of the growing season (summer–autumn), as a result of size-dependent emigration, length–frequency data are biased by the decline in abundance of larger individuals, causing an apparent reduction in growth rates. Negative daily growth rates were not considered. At the end, only those data sets with >5 growth rate estimates were used in subsequent analyses.

**DEB model-based growth potential**

Growth potential was determined based on the standard DEB model (Kooijman 2010). A thorough description of the model and relevant equations is given in Van der Veer et al. (2010; Eqs. 1–3). Maximum food conditions were simulated in the model using the scaled functional response (f), a dimensionless parameter ranging from 0 to 1 (ad libitum food) that relates ingestion to food density in the environment through a Holling type II functional response (Kooijman 2010). Under constant food conditions (or at abundant food due to the hyperbolic functional response), DEB differential equations can be analytically solved and the dynamics of growth then simplifies to Von Bertalanffy growth (Kooijman 2010).

Growth rates are affected not only by food availability but also by temperature conditions, due to the temperature dependence of physiological rates. In the DEB model, such thermal dependence is described using an Arrhenius-type relationship extended to account for the effects of temperature at both lower and upper tolerance boundaries (Kooijman 2010, p. 21; Van der Veer et al. 2010, Eq. 4).

**Realized growth ratio**

Field growth rates were compared with the DEB-predicted growth potential to evaluate the possibility of food limitation. Growth potential, defined as the

<table>
<thead>
<tr>
<th>Location</th>
<th>Latitude</th>
<th>Species</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valosen, Norway</td>
<td>67° N</td>
<td><em>P. platessa</em></td>
<td>Freitas et al. (2010a)</td>
</tr>
<tr>
<td>Faxaflói¹, SW Iceland</td>
<td>64° N</td>
<td><em>P. platessa</em></td>
<td>Hjörleifsson &amp; Pálsson (2001)</td>
</tr>
<tr>
<td>Gullmar Fjord², Sweden</td>
<td>58° N</td>
<td><em>P. platessa</em></td>
<td>L. Pihl unpubl. data</td>
</tr>
<tr>
<td>Loch Ewe, Scotland</td>
<td>57°50’N</td>
<td><em>P. platessa</em></td>
<td>Steele &amp; Edwards (1970), Edwards &amp; Steele (1968)</td>
</tr>
<tr>
<td>Mariager Fjord³, Denmark</td>
<td>56° N</td>
<td><em>P. flesus</em></td>
<td>Andersen et al. (2005)</td>
</tr>
<tr>
<td>Port Erin Bay, Isle of Man, UK</td>
<td>54° N</td>
<td><em>P. platessa</em></td>
<td>R. Nash unpubl. data</td>
</tr>
<tr>
<td>Dollard estuary, eastern Dutch Wadden Sea</td>
<td>53° N</td>
<td><em>P. platessa</em></td>
<td>Jager et al. (1995)</td>
</tr>
<tr>
<td>Balgzand, western Dutch Wadden Sea</td>
<td>53° N</td>
<td><em>P. platessa</em>, <em>S. solea</em></td>
<td>Van der Veer et al. (2010), present study</td>
</tr>
<tr>
<td>Minho Estuary, Portugal</td>
<td>41° N</td>
<td><em>P. flesus</em></td>
<td>Dolbeth et al. (2010), present study</td>
</tr>
<tr>
<td>Mondego Estuary, Portugal</td>
<td>40° N</td>
<td><em>P. flesus</em></td>
<td>Martinho et al. (2008), Dolbeth et al. (2010)</td>
</tr>
<tr>
<td>Tagus Estuary⁴, Portugal</td>
<td>39° N</td>
<td><em>S. solea</em></td>
<td>Martinho et al. (2008), Dolbeth et al. (2010)</td>
</tr>
</tbody>
</table>

Table 1. *Pleuronectes platessa*, *Platichthys flesus* and *Solea solea*. Locations and original sources of the growth data of plaice flounder and sole used in the present analyses. Superscripts indicate additional temperature sources: ¹Marine Research Institute (Station Reikjavik; www.hafro.is/Sjora); ²Swedish Meteorological and Hydrological Institute (Station Alsbäck, Slåggö; www.smhi.se); ³National Environmental Research Institute (MADS; Station Mariager Fjord, Dybet; www.dmu.dk/en/water/marinemonitoring/mads); ⁴Cabral & Costa (2001).
maximum growth rate at constant and maximum food conditions ($f = 1$), was determined for each growth interval using site-specific water temperatures.

DEB parameters were taken from Van der Veer et al. (2001b, 2009, 2010) and Freitas et al. (2010b). Before any simulations, the reliability of the model parameter values was validated by comparing model-predicted maximum growth rates with laboratory-derived maximum growth. For plaice and flounder, experimental growth data by Fonds et al. (1992) were used, while for sole, data were taken from Fonds & Rijnsdorp (1988) and Irvin (1973 in Howell 1997).

Temperature parameters used to correct growth rates according to the extended Arrhenius relationship were also based on previous estimates (Freitas et al. 2010b) and in some further fine-tuning by comparing the shape of the growth curve predicted by the DEB model with the shape of those derived from experimental observations. For sole, Arrhenius temperature was changed from 8500 to 6500 K to better match the growth curves described by the 2 data sets available. Although growth rates described in Irvin’s experiments are lower than those in Fonds & Rijnsdorp (1988), both data sets show little increase in growth rates above 20–22°C.

Due to gender-related differences in energy assimilation, some of the parameters differ (Freitas et al. 2010b), therefore, maximum growth in relation to temperature was predicted for each sex separately. However, for comparison with field growth where sexes were not distinguished, DEB-predicted growth rates for females and males were averaged, assuming a 1:1 field sex ratio.

The fraction of maximum growth realized by juvenile fish of each species was estimated as the ratio between observed and DEB-predicted maximum growth (realized growth ratio [RG], dimensionless). Patterns in RG were analyzed over the growing season and compared among areas to assess trends with latitude. To help visualizing patterns in data, smoothing curves were added using the LOESS function in R (R Development Core Team 2009) and the fitted values in each month were plotted against latitude for each species.

**RESULTS**

**Field growth**

In total, reliable information could be collected from 11 areas ranging from northern Norway to the Portuguese coast (Table 1). In most cases, only studies of single species were available. Time coverage of the data sets varied from single to multiple years. Since interannual comparisons were outside the scope of this paper, data from different years were pooled.

Observed field growth rates showed high variability in all areas in the 3 species (Fig. 1). Flounder and plaice growth rates were mostly up to 0.10 cm d$^{-1}$, with the exception of a few individual data points in the order of 0.13 cm d$^{-1}$ for plaice in Swedish waters (Fig. 1a). In sole, maximum growth rates were between 0.15 and 0.20 cm d$^{-1}$ reflecting higher growth potential compared with the other species (Fig. 1c). In all species, growth rates seem to decline...
throughout the season with the exception of Mondego Estuary (southern Europe), where autumn growth rates in sole were still considerably high.

### DEB predictions and RG

DEB parameters are summarized in Table 2. As expected, maximum growth rates predicted by the DEB model differed among sexes, with higher values for females. Overall, DEB parameter sets appeared to be reliable, as the resulting growth rates at various temperatures were in the same order of magnitude from those established in the laboratory and reproduced a similar temperature effect (Fig. 2). For plaice and flounder, DEB simulations for males were close to experimental growth, but for females, predictions were higher, particularly in plaice (Fig. 2).

In general, RG decreased over time, indicating an increasing discrepancy between field growth and model predictions. For plaice, most areas had similar patterns with relatively higher values in the beginning of the season, followed by a continuous decrease over time or stabilization, as observed in the Balgzand (Fig. 3a). The only exception was France, where RG seemed to increase over the course of time. Similar temporal dynamics were observed in

Table 2. *Pleuronectes platessa*, *Platichthys flesus* and *Solea solea*. Dynamic energy budget (DEB) model parameters at a species-specific reference temperature, and temperature dependence parameters for the 3 flatfish species. Whenever parameters differ between sexes, values for males are shown in (). For details on the notation used see Kooijman (2010)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>Interpretation</th>
<th>Plaice</th>
<th>Flounder</th>
<th>Sole</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_1$</td>
<td>K</td>
<td>Reference temperature</td>
<td>283</td>
<td>283</td>
<td>293</td>
</tr>
<tr>
<td>$v$</td>
<td>cm d$^{-1}$</td>
<td>Energy conductance</td>
<td>0.156</td>
<td>0.156</td>
<td>0.435</td>
</tr>
<tr>
<td>$g$</td>
<td>–</td>
<td>Energy investment ratio</td>
<td>2.635 (3.930)</td>
<td>3.590 (4.211)</td>
<td>3.111 (3.684)</td>
</tr>
<tr>
<td>$k_M$</td>
<td>d$^{-1}$</td>
<td>Maintenance rate coefficient</td>
<td>0.0035</td>
<td>0.0035</td>
<td>0.0097</td>
</tr>
<tr>
<td>$L_m$</td>
<td>cm</td>
<td>Maximum physical length</td>
<td>78 (52)</td>
<td>56 (47)</td>
<td>75 (64)</td>
</tr>
<tr>
<td>$\delta_M$</td>
<td>–</td>
<td>Shape coefficient</td>
<td>0.219</td>
<td>0.224</td>
<td>0.192</td>
</tr>
<tr>
<td>$T_A$</td>
<td>K</td>
<td>Arrhenius temperature</td>
<td>7000</td>
<td>7500</td>
<td>6500</td>
</tr>
<tr>
<td>$T_L$</td>
<td>K</td>
<td>Lower boundary of tolerance range</td>
<td>277</td>
<td>296</td>
<td>301</td>
</tr>
<tr>
<td>$T_H$</td>
<td>K</td>
<td>Upper boundary of tolerance range</td>
<td>295</td>
<td>296</td>
<td>301</td>
</tr>
<tr>
<td>$T_{AL}$</td>
<td>K</td>
<td>Rate of decrease at lower boundary</td>
<td>50 000</td>
<td>35 000</td>
<td>35 000</td>
</tr>
<tr>
<td>$T_{AH}$</td>
<td>K</td>
<td>Rate of decrease at upper boundary</td>
<td>75 000</td>
<td>75 000</td>
<td>50 000</td>
</tr>
</tbody>
</table>

**Fig. 2.** *Pleuronectes platessa*, *Platichthys flesus* and *Solea solea*. Maximum daily growth of juvenile (a) plaice, (b) flounder and (c) sole under optimal food conditions in relation to water temperature. Growth predictions based on laboratory-derived models are indicated by filled circles (in a,b: ● = Fonds et al. 1992; in c: ● = Fonds & Rijnsdorp 1988, ● = Irvin 1973 in Howell 1997). Lines: DEB model predictions for females (solid) and males (dashed)
flounder (Fig. 3b), with steep declines in RG over time, followed by stabilization at around the end of July. In Balgzand, no clear change was observed with time and field growth rates were ~70% (RG = 0.7) of the potential growth. Sole showed a more variable pattern compared with the other 2 species (Fig. 3C). Tagus Estuary was the only site with a clear and continuous decrease in RG over time. In the remaining areas, dome-shaped patterns were observed, with higher differences in observed relative to potential growth rates at both the beginning and end of the season. In Mondego Estuary, there was an additional increase in RG in early autumn.

Latitudinal comparisons of RG were made for each month (May to September) using the predicted values from the smooth curves (Fig. 4). The RG of plaice and flounder exhibited clear positive trends with latitude from June to August. In September, values were similar among locations, ranging from 20 to 40% of maximum potential growth. In sole, the same tendency was observed during July and August, with higher RG in northern compared with southern nurseries. In June, however, the trend was reversed, indicating close to maximum field growth rates in southern areas and lower than expected growth at higher latitudes (Eastern Channel and Balgzand).

Differences among nurseries were highest in June in plaice and flounder, as shown by steep regression lines that progressively decreased towards the end of the growing season (Fig. 4). In sole, differences between areas were largest in July.

**DISCUSSION**

**Habitat quality of flatfish nursery areas**

Shallow soft-bottom areas are assumed to provide abundant food supply to epibenthic assemblages (Evans 1983). This conjecture is supported by an enhanced diversity and abundance of macrobenthic fauna in flatfish nursery grounds compared with adjacent non-nursery areas (Wouters & Cabral 2009).

Using a mechanistic approach for capturing growth dynamics, the present study challenges the assumption of stability in food conditions and supports previous indications that habitat quality in terms of growth conditions changes over the season, not only for plaice but also flounder, and that this pattern is common to several nurseries. Similar temporal dynamics, with a progressive deterioration of growth conditions throughout summer, have been observed for juvenile plaice in Tralee Beach and Loch Caolisport Beach, western Scotland, using an RNA-based growth index (Ciotti et al. 2010). The agreement in results using different methods seems to support a widespread phenomenon of food limitation.
Available resources can be affected by exploitative competition between individuals, either of the same or different species. Studies with plaice point to a lack of intraspecific competition for food (Van der Veer et al. 2010) and for Irish Sea populations, it is even suggested that the carrying capacity of nursery grounds for plaice is rarely approached (Nash et al. 2007). However, interactions within the benthic-feeder assemblage (other fish and invertebrates) can modify the carrying capacity and trajectories for individual species (Gibson 1994, Nash et al. 2007). An analysis similar to the one applied in this study has suggested that in sand gobies Pomatoschistus minutus occurring sympatriacally with juvenile flatfish during summer, growth is close to the maximum possible across their distribution range (Freitas et al. 2011), suggesting a competitive advantage.

Maximum individual intake rates can also be reduced as a consequence of interference by other organisms, affecting prey searching, capturing or handling by flatfish (see Van der Veer et al. 2010). Predator presence, for instance, has been found to reduce feeding activity in plaice (Burrows & Gibson 1995) and sole (Maia et al. 2009). Another example of interference competition reducing the amount of food available for juvenile flatfish is the browsing activity by other fish and crustaceans on bivalve and polychaete soft tissues (De Vlas 1979, 1985, Kamermans & Huitema 1994), affecting prey quality (Bergman et al. 1988a, Zwarts & Wanink 1989, Zajac 1995, De Goeij et al. 2001, Meyer & Byers 2005).

The temporal trend in growth conditions observed in plaice and flounder was not exhibited by sole, which seems to suggest that even if the underlying mechanism is the same, responses cannot be generalized even within a functional group. Although the reasons are not clear, sole may avoid interspecific competition by spatial and temporal feeding activities (Besyst et al. 1999), for instance, taking advantage of its capacity to forage during the night as opposed to plaice and flounder, which are daytime feeders (Lagardère 1987, Besyst et al. 1999).

Our approach to analyse changes in habitat quality based on RG relies on the premise of accurate characterization of both field growth and temperature conditions experienced. This study builds on previous research, taking advantage of available data, which invariably describe growth from changes in modal or mean length of sampled populations rather than from individual trajectories. These estimates are known to be influenced by migration, gear efficiency and size-selective mortality. Although ideally one could avoid some bias associated with migrations by considering only the period once settlement is completed and before the autumn migrations to deeper waters start, this time window is probably nursery-ground-dependent (Nash et al. 2007), making the comparison between areas more complicated. Therefore, a more general criterion for data selection had to be used. By restricting the analyses to a period where a more or less linear increase in size was observed, we have tried to preclude inaccurate estimations, which would affect the evaluation of realized growth.

Some of the discrepancy between field and maximum predicted growth rates can be related to an inaccurate representation of field temperature conditions. As a result of fine-scale habitat selection among juveniles, the thermal regime experienced by the population may differ from the temperature recorded in the field (Hurst et al. 2010). In some cases,
this could result in RG values larger than 1 if, for instance, juveniles tend to select warmer locations early in the season (Fig. 3).

**Latitudinal trends**

The temporal dynamics in realized growth within each area complicates a latitudinal comparison of habitat quality for the various species. Nevertheless, our results seem to indicate a consistent positive latitudinal trend during summer in all 3 species studied. Despite the suggestion for lower benthic biomass and production at lower latitudes (Pauly 1994), evidence from a few European nurseries does not seem to support such a trend, instead indicating similar biomass figures (Beukema et al. 2002, Wouters & Cabral 2009, Freitas et al. 2010a). From a bioenergetics viewpoint, assuming that food conditions are generally similar, energy uptake must be higher in order to compensate for extra metabolic costs at southern, warmer locations. In the presence of competition, southern areas are thus more prone to food limitation, as individual energy intake is likely to be reduced compared with more northern areas.

We have observed a downward shift in RG patterns from northern to southern areas, which supports this competition hypothesis. The results for sole are particularly indicative of how this mechanism may act: in the beginning of the growing season, the RG of sole is higher in southern compared with northern areas; however, throughout the season this pattern inverts. In contrast to northern European areas where sole only occur after plaice and flounder have settled (Jager et al. 1995, Amara et al. 2001, Van der Veer et al. 2001a), in southern European nurseries, sole juveniles are the first marine species to colonize these areas, appearing in April–May (Cabra et al. 2003, Dolbeth et al. 2010), and in some years, young-of-the-year can even be observed from late January onwards (Martinho et al. 2008). This could confer an advantage to sole by occupying an empty niche with ample food resources, which, combined with high summer temperatures, would allow for fast growth. Throughout the season, competition starts playing its role in reducing food availability and leading to the same spatial trends as observed in flounder and plaice.

Southern European nursery areas do not allow for maximum growth, neither in flounder nor in sole, in spite of the longer growing seasons which lead to larger juveniles by the end of the first year (Martinho et al. 2008). In the Tagus Estuary, food has not been considered a limiting factor for sole based on comparisons of food consumption and prey biomass (Vinagre & Cabral 2008). Instead, thermal stress has been suggested as a possible cause for hampering growth rates in sole as summer temperatures largely exceed metabolic optimum temperatures (Vinagre et al. 2008). Excessive temperatures have also been suggested as the mechanism responsible for the drastic decline in flounder observed in recent years in this estuary (Cabra et al. 2001, 2007), and for a disruption in plaice recruitment in the Bay of Biscay (Désaunay et al. 2006, Hermant et al. 2010).

In the context of global warming, our results suggest that habitat quality of southern European nurseries for juvenile growth may be particularly affected by climatic changes through a combination of both food and thermal constraints. Our approach explicitly accounted for the thermal effects on growth rates. However, the parameters used to describe temperature dependence were taken as constant on both temporal and spatial scales. Temperature has such a pervasive effect on performance that organisms have certainly managed to respond to their thermal environment. These responses can vary from short-term organismal-level behavioural and physiological adjustments, to adaptive shifts within the lifetime of an individual (Angilleta et al. 2002) and, on the scale of multiple generations, genetic changes may occur (Braddock & Holzapfel 2006). Differences in growth performance among flatfish populations and some evidence for latitudinal compensation in growth have already been demonstrated (Jmsland et al. 2000, 2001, Jonassen et al. 2000). However, how much of this intraspecific variability translates into the energy budget and which model parameters are more likely to be affected still need to be clarified. Elucidating these aspects will bring us one step forward towards better predicting the responses in juvenile nursery quality to changes in climate.

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LITERATURE CITED

- Gibson RN, Robb L, Wennhage H, Burrows MT (2002) Ontogenetic changes in depth distribution of juvenile flat-
floses in relation to predation risk and temperature on a
shallow-water nursery ground. Mar Ecol Prog Ser 229:
233–244

Henderson PA, SeabY BM, Somes JR (2006) A 25-year study of
climatic and density-dependent population regulation of
common shrimp Crangon crangon (Crustacea: Car
idea) in the Bristol Channel. J Mar Biol Assoc UK 86:
287–298

Hermant M, Lobry J, Bonhommeau S, Poulard JC, Le Pape
O (2010) Impact of warming on abundance and occurrence
of flatfish populations in the Bay of Biscay (France).
J Sea Res 64: 45–53

Hjörleifsson E, Pálsson J (2001) Settlement, growth and mor
tality of 0-group plaice (Pleuronectes platessa) in Ice

Howell BR (1997) A re-appraisal of the potential of the sole,
Solea solea (L.), for commercial cultivation. Aquaculture
155: 355–365

Hurst TP, Abookire AA (2006) Temporal and spatial varia
tion in potential and realized growth rates of age-0 year

Hurst TP, Abookire AA, Knott B (2010) Quantifying thermal
effects on contemporary growth variability to predict
responses to climate change in northern rock sole (Lepi

Imsland AK, Foss A, Nævdal G, Cross T, Bonga SW, van
Ham E, Stefansson SO (2000) Countergradient variation in
growth and food conversion efficiency of juvenile tur
bot. J Fish Biol 57: 1213–1226

Imsland AK, Foss A, Stefansson SO (2001) Variation in food
intake, food conversion efficiency and growth of juvenile
turbot from different geographic strains. J Fish Biol 59:
449–454

0-group flatfish in the brackish Dollard (Ems estuary,

Jonassen TM, Imsland AK, FitzGerald R, Stefansson MÖ
and others (2000) Geographic variation in growth and
growth efficiency of juvenile Atlantic halibut related to
latitude. J Fish Biol 56: 279–294

Kamermans P, Huitema HJ (1994) Shrimp (Crangon crangon
L.) browsing upon siphon tips inhibits feeding and
growth in the bivalve Macoma balitica (L.). J Exp Mar
Biol Ecol 175: 39–75

Kooijman SALT (2000) Dynamic energy and mass budgets in
biological systems. Cambridge University Press, Cam
bridge

Kooijman SALT (2010) Dynamic energy budget theory for
metabolic organization. Cambridge University Press, Cam
bridge

Lagardère JP (1987) Feeding ecology and daily food con
sumption of common sole, Solea vulgaris Quensel, juve

of coastal and estuarine essential fish habitats: estima
tions based on the size of juvenile common sole (Solea
solea L.). Estuar Coast Shelf Sci 58: 793–803

Maes J, Taillieu A, Van Damme PA, Cottenie K, Ollevier FP
(1998) Seasonal patterns in the fish and crustacean com
munity of a turbid temperate estuary (Zeeschelde Estu
ary, Belgium). Estuar Coast Shelf Sci 47: 143–151

the foraging behaviour of Solea senegalensis. J Mar
Biol Assoc UK 89: 645–649

Martinho F, Leiiao R, Neto JM, Cabral H, Lagardere F,
Pardal MA (2008) Estuarine colonization, population struc
ture and nursery functioning for 0-group sea bass (Dicen
trarchus labrax), flounder (Platichthys flesus) and sole (Solea
solea) in a mesotidal temperate estuary. J Appl Ichthyol
24: 229–237

Meyer JJ, Byers JE (2005) As good as dead? Sub lethal pre
dation facilitates lethal predation on an intertidal clam.
Ecol Lett 8: 160–166

of juvenile plaice (Pleuronectes platessa L.) on a small
Irish Sea nursery ground (Port Erin Bay, Isle of Man, UK).

Dynamics of shallow-water juvenile flatfish nursery
grounds: application of the self-thinning rule. Mar Ecol
Prog Ser 344: 231–244

ot ML, Hureau JC, Nielsen J, Tortonec E (eds) Fishes of
the North-eastern Atlantic and Mediterranean. United
Nations Educational, Scientific and Cultural Organiza
tion, Paris, p 1299–1307

Pauly D (1994) A framework for latitudinal comparisons of

Pihl L (1985) Food selection and consumption of mobile
epibenthic fauna in shallow marine areas. Mar Ecol Prog
Ser 22: 169–179

Quéro JC, Desoumter M, Lagardère F (1986) Soleidae. In:
Whitehead PJP, Bautch ML, Hureau JC, Nielsen J, Tor
tonec E (eds) Fishes of the North-eastern Atlantic and
Mediterranean. United Nations Educational, Scientific
and Cultural Organization, Paris, p 1308–1324

R Development Core Team (2011) R: A language and envi
ronment for statistical computing. R Foundation for Sta
tistical Computing, Vienna, Austria. www.R-project.org

Rijndorp AD, Van Beek FA, Flatman S, Millner RM, Riley
JD, Giret M, De Clerck R (1992) Recruitment of sole
stocks, Solea solea (L.), in the Northeast Atlantic. Neth J

tion regulation and speciation. University of Washing
ton Press, Seattle, WA

Sogard SM (1997) Size-selective mortality in the juvenile stage

plaice and common dabs in Loch Ewe, IV. Dynamics of
4: 174–187

Teal LR, De Leeuw JJ, Van der Veer HW, Rijndorp AD
(2008) Effects of climate change on growth of

Van der Veer HW (1986) Immigration, settlement, and den
sity-dependent mortality of a larval and early postlarval
0-group plaice (Pleuronectes platessa) population in the

Van der Veer HW, Bergman MJN, Dapper R, Witte JJ
(1991) Population dynamics of an intertidal 0-group
flounder Platichthys flesus population in the western

Van der Veer HW, Berghahn R, Rijndorp AD (1994) Impact
of juvenile growth on recruitment in flatfish. Neth J Sea
Res 32: 153–173

Van der Veer HW, Berghahn R, Miller JM, Rijndorp AD
(2000) Recruitment in flatfish, with special emphasis on
North Atlantic species: progress made by the Flatfish
Fig. A1. Average monthly water temperatures (●) recorded at the various locations, compiled from the original published studies or oceanographic data centres. For data sources, see Table 1. ○: temperature values used to estimate maximum growth rates.