

Physiology and Biogeography: The Response of European Mussels (Mytilus spp.) to Climate Change *

Authors: Fly, Elizabeth K., Hilbish, Thomas J., Wethey, David S., and Rognstad, Rhiannon L.

Source: American Malacological Bulletin, 33(1): 136-149

Published By: American Malacological Society

URL: https://doi.org/10.4003/006.033.0111

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at www.bioone.org/terms-of-use.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Physiology and biogeography: The response of European mussels (*Mytilus* spp.) to climate change*

Elizabeth K. Fly^{1,2}, Thomas J. Hilbish³, David S. Wethey³, and Rhiannon L. Rognstad^{3,4}

¹South Carolina Sea Grant Consortium, Charleston, South Carolina, 29401, U.S.A.

²Carolinas Integrated Sciences and Assessments, Department of Geography, University of South Carolina, Columbia, South Carolina 29208, U.S.A.

Correspondence, Elizabeth Fly: lizkfly@gmail.com

Abstract: To understand how ecological communities may respond to climate change we have adopted the approach of determining the response of major ecosystem engineers that determine community composition and function. We utilize two approaches, correlative and mechanistic, to understand the current and future distributions of the marine mussels *Mytilus edulis* Linnaeus, 1758 and *M. galloprovincialis* Lamarck, 1819 in Europe. Both are dominant space-occupying species that control biodiversity in many coastal ecosystems and are the basis of the largest aquaculture production in Europe. A mechanistic analysis of physiological energetic response to temperature of the two species indicates that *M. edulis* cannot sustain a positive energy balance for sustained periods when sea surface temperature (SST) is greater than 23 °C, while *M. galloprovincialis* can maintain a positive energy balance at SST up to 30 °C. There is no difference in energetic response of the two species at cold temperatures (5–10 °C). The upper temperature threshold of positive energy balance in each species corresponds closely to the distribution of SST at their respective southern range limits in Europe. Alternatively, the northern range limit of *M. galloprovincialis* coincides with areas where winter SST is less than 9 °C, but there is no evidence of an energetic limit to this species at the cold end of its geographic range. Presently there is no mechanistic explanation for the difference between species in their northern range limits; however, as indicated by Random Forest modeling, *M. galloprovincialis* appears to be limited by cold temperatures during winter, suggesting the hypothesis of failure in reproductive development. These approaches allow for the ability to forecast changes in the distributions of these two species in Europe as SST continues to increase.

Key words: Species distribution modeling, Random Forest, blue mussel, ecological forecasting

It is now well established that organisms are responding to the unprecedented rate of climate warming, including changes in the timing of biological activities (Philippart et al. 2003, Edwards and Richardson 2004, Burrows et al. 2011) and biogeography (Burrows et al. 2011, Hilbish et al. 2012, Jones et al. 2012). To understand current and potential distributions of species, many biologists rely on species distribution models (SDMs) that use spatial environmental data relevant to a species to predict its distribution (Guisan and Thuiller 2005, Elith and Leathwick 2009, Peterson et al. 2011). There are two types of SDMs: correlative and mechanistic, though the two are not mutually exclusive. Correlative SDMs create predictions based on correlations between species distribution records and environmental variables (Robertson et al. 2003, Buckley et al. 2010). These types of models can be generated fairly quickly and can often, particularly for a species for which little is known, provide more information on

what specific environmental variables may be important for the distribution of that species (Robertson *et al.* 2003, Elith and Leathwick 2009). Mechanistic SDMs create predictions based on knowledge of underlying physiological and/or life history responses to environmental variables (Robertson *et al.* 2003, Kearney and Porter 2009, Buckley *et al.* 2010, Woodin *et al.* 2013). Mechanistic SDMs are more time-consuming to develop, since the important physiological mechanisms that potentially regulate the species distribution must first be determined (Kearney and Porter 2009), but may provide greater insight into the factors limiting a species' distribution than a purely correlative model.

The current challenge is to move beyond modeling changes in the range limits of single species to develop an understanding of the biological mechanisms responsible for these changes and, importantly, the responses of communities and the functions of ecosystems. We have taken the

³Department of Biological Sciences, University of South Carolina, Columbia, South Carolina 29208, U.S.A.

⁴Present address: Keck Science Department, Claremont McKenna College, Claremont, California, 91711, U.S.A.

^{*}Symposium on "Climate Change and Molluscan Ecophysiology" at the 79th Annual Meeting of the American Malacological Society, 2013 World Congress of Malacology, July 23, 2013 Azores.

approach of focusing on the response of key ecosystem engineers that both control the composition and diversity of ecological communities and are primary determinants of the "function" of ecosystems with respect to the goods and services provided to human societies, using both correlative and mechanistic methods. Mechanistic SDMs have been suggested as a more responsive model to climate change and invasive species scenarios (Kearney and Porter 2009) as they are better able to extrapolate to novel environmental conditions due to an understanding of the causative mechanisms behind species distributions (Helmuth et al. 2005, Kearney and Porter 2009, Woodin et al. 2013). However, these two types of approaches are not mutually exclusive and we show that incorporating both may provide a more robust model of a species' complete biogeographic range (Lawler et al. 2006, Elith and Leathwick 2009, Sará et al. 2011).

In this study our focal system is marine mussels in the genus Mytilus across Linnaeus, 1758 a European scale. We selected mussels because they are major determinants of space and, therefore, biodiversity within rocky intertidal marine communities (Suchanek 1992). Also, mussels are farmed commercially, and mussel aquaculture is a multibillion dollar industry (FAO 2012). Mytilus edulis Linnaeus, 1758 and M. galloprovincialis Lamarck, 1819 are sister species that inhabit the coastal waters of Europe. Mytilus edulis is a cold-temperate species, inhabiting northern European waters to the French/Spanish border in the Bay of Biscay (Hilbish et al. 2012). Mytilus galloprovincialis is native to the Mediterranean and is found as far north as the British Isles (Skibinski et al. 1983, McDonald et al. 1991, Seed 1992). The overlap of these species has created a mosaic hybrid zone, with alternating patches of hybrids and pure parental populations from the Bay of Biscay to northern Scotland (Skibinski et al. 1983, Coustau et al. 1991, Bierne et al. 2003, Gosling et al. 2008, Hilbish et al. 2012). We sought to understand what controls the biogeographic distributions of these economically and ecologically important species in Europe, and how their functional performance differs within their ranges. This work provides a basis for forecasting future changes in the distribution of these species as the climate continues to change.

MATERIALS AND METHODS

European *Mytilus* distributions

Mytilus spp. have been well studied for many decades in Europe (Skibinski et al. 1983, Coustau et al. 1991, Gardner and Skibinski 1991, Rawson and Hilbish 1998, Daguin et al. 2001, Bierne et al. 2003), and their distribution in the past decade in various regions has been documented quite thoroughly. Much of this distribution was reported in Hilbish et al.

(2012), who sampled 97 locations along the Atlantic coast of Spain and France. However, we developed an even more extensive range database by collating data from several studies (Lionetto et al. 2003, Gosling et al. 2008, Pisanelli et al. 2009, Banni et al. 2011, Kijewski et al. 2011, Hilbish et al. 2012, Wethey unpubl. data, Appendix A) as well as sampling in certain regions that were missing recent data (particularly the United Kingdom). Although these species do hybridize, for the purposes of this study we focused on "pure" species populations (populations that contained ≥ 95% allele frequency of one species) because the physiological studies of Fly and Hilbish (2013), described below, focused on "pure" species rather than hybrids. In 2008 and 2009, we sampled mussel populations at 43 sites in England, Scotland, and Wales that were originally sampled in 1976 and 1977 by Skibinski et al. (1983). To gain higher spatial resolution of the hybrid zone in southwest England, we sampled an additional 24 sites in 2010 that were previously sampled in 1996 by Hilbish et al. (2002). DNA extraction and genotyping at the Glu-5' locus followed the protocols in Hilbish et al. (2012).

Modeling

We used two separate, yet complementary, modeling approaches to identify key variables determining the range limits of *Mytilus edulis* and *M. galloprovincialis*. The correlative approach was used primarily to classify known sites of "pure" populations of one of the two species, and then identify temperature variables that could be used to differentiate between *M. edulis* sites and *M. galloprovincialis* sites. The mechanistic approach used laboratory estimates of scope for growth of the two species to compare regions where scope for growth is estimated to be zero with the geographic limits of the two species.

Correlative approach

To estimate the importance of sea surface temperature (SST) on the distribution of *Mytilus* spp. in Europe, we analyzed the distribution of "pure" Mytilus populations using Random Forest (RF) modeling, which is becoming increasingly popular in ecological studies (Cutler et al. 2007). This machine learning model generates a summary of many classification trees to determine the best predictor variables for a dataset (Breiman 2001). To build the model, we classified sites from both our field studies and the literature (see Appendix A) as either M. galloprovincialis or M. edulis based on allele frequency with a 95% threshold; sites with ≥ 95% of alleles for a particular species were classified as a population of that species. We used the temperature parameters of yearly mean SST, monthly mean SST (for each month), and seasonal mean SST, as in Hilbish et al. (2012). These parameters were calculated using National Oceanic and Atmospheric Administration (NOAA) Optimum Interpolation 1/4 Degree

Daily SST (OISST) data derived from Advanced Very High Resolution Radiometer (AVHRR) satellite sensors (Reynolds et al. 2007), acquired from the National Climatic Data Center archive at http://www.ncdc.noaa.gov/thredds/catalog/oisst/NetCDF/AVHRR/catalog.html. The SST value for each sample site was estimated by finding the pixel on each SST grid nearest to the sampling location. We determined SST means listed above for each population for the five years prior to each mussel sample date. This time period of five years was chosen to encompass on average the life span of the mussels (Hilbish et al. 2012).

The RF model was built using the randomForest package (Liaw and Wiener 2002) in R version 2.7.1 (R Development Core Team 2010) with 250,000 individual trees, in which one-third of the cases were left out of each tree to be used as an estimate of classification error as the trees were added to the forest. Classification error is the frequency of test cases (those not included in model generation) incorrectly classified by the final model and was used to evaluate the ability of the model to differentiate between M. galloprovincialis and M. edulis sites. The relative importance of each temperature variable was determined for the RF model to identify which variables were most useful for classifying the sites, thus, identifying temperature differences between the ranges of the two species. The importance of the predictor temperature variables was determined using the mean decrease in accuracy calculated during the RF analysis. This metric is calculated by identifying the decrease in the accuracy of the model when the values of a single predictor variable are randomly permuted. Thus, variables with the larger mean decrease in accuracy are more important for data classification.

Mechanistic approach

The physiological energetics of Mytilus edulis and M. galloprovincialis were measured by Fly and Hilbish (2013). They integrated measurements of filtration rate, absorption efficiency, and metabolic rate to derive an estimate of the scope for growth (SFG) (sensu Widdows and Bayne 1971 the SFG measures the energy above maintenance available for growth and reproduction) of mussels at water temperatures ranging from 5 °C to 30 °C. We used their estimate of the critical SST at which SFG = 0 to determine the geographic regions in Europe in which each species can potentially maintain a permanent population. In regions that exceed the critical SST at which SFG = 0, populations of each species are expected to be absent or ephemeral (Sokolova et al. 2012, Woodin et al. 2013). We compared the distribution of *M. edulis* and *M. gal*loprovincialis to the monthly SST along the European coastline and to the critical temperature at which SFG = 0 (Fly and Hilbish 2013). SFG data were collected from one population of each species (Fly and Hilbish 2013). While local adaptation may be possible, Mytilus populations have the potential for very high gene flow, with dispersal distances of up to 100 km (Gilg and Hilbish 2003). The data presented below suggest that there is not local adaptation occurring in genes relevant to setting *Mytilus* range limits, since the physiological energetics measured using animals from one population appear to well explain the southern range edge of both *M. edulis* and *M. galloprovincialis*. If there were local adaptation occurring, we would expect to see these data fail to explain distribution limits.

Ecological forecasting

Forecasts of future environmental conditions were made using model output from the EURO-CORDEX project (Jacob et al. 2014). This project uses regional circulation models (RCM) to downscale global circulation models (GCM) in order to predict monthly average SST on a 44 km grid. We chose to use the 4.5 Wm⁻² (rcp4.5) global energy imbalance scenario of the IPCC 5th Assessment Report, which simulates moderate future warming and is similar to the IPCC 4th Assessment Report scenario A1B. To date, 8 GCMs have been downscaled with one RCM (Appendix B), and the results have been posted on the Earth System Grid (http://esgf-data. dkrz.de/esgf-web-fe/). EURO-CORDEX 44-km data were on a curvilinear grid, so they were interpolated to a 25 km Mercator grid, using the R package "akima" (Akima et al. 2013). Since there is disagreement among the models in terms of the spatial distribution of sea surface temperatures, we made maps of the fraction of models that predicted SST of at least 20 °C in summer (August) and at least 9 °C in winter (February). The temperature of 20 °C in summer is a proxy for the southern geographic limit of M. edulis (Fly and Hilbish 2013) and the temperature of 9 °C is a proxy for the northern geographic limit of M. galloprovincialis (Hilbish et al. 2012), as indicated by the mechanistic and correlative models. We used the median of the model predictions as our estimate of the expected geographic distribution of the Mytilus species over the periods 2046–2050 and 2096–2100. For comparison to the present, we used the daily NOAA OISST product (Reynolds et al. 2007) to make maps of monthly average SST over the period 2002-2009. We then determined for each pixel in the map, the fraction of years (2002–2009) with temperatures higher than 9 °C in winter and 20 °C in summer.

RESULTS

The distribution of *Mytilus* spp. in Europe ranges from northern Africa, through the Mediterranean, along the Atlantic coastline into Scandinavia and on the British Isles (Fig. 1). Contiguous populations of *M. galloprovincialis* extend from the Mediterranean north to the border between Spain and France in the Bay of Biscay (Fig. 1). Two disjunct

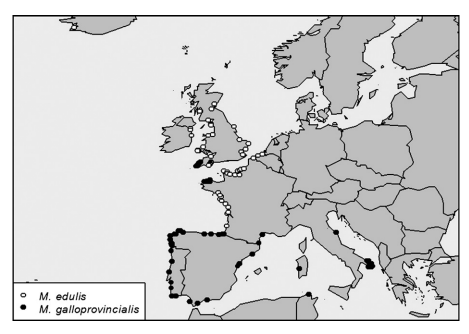


Figure 1. Sample locations used for biogeographic modeling of *Mytilus* spp. in Europe. These data were collated from several published studies (see Appendix A) as well as additional sampling in certain regions that were missing data (in particular the United Kingdom). For the purposes of this study we focused on "pure" species populations, and we defined "pure" as populations that contained \geq 95% genotype of either *M. edulis* or *M. galloprovincialis*. Although the data points are sparse in the Mediterranean and North Sea, these areas are nearly exclusively *M. galloprovincialis* and *M. edulis*, respectively. The current southern limit of *M. edulis* occurs at the France/Spain border in the Bay of Biscay, while the current northern limit of *M. galloprovincialis* is on France's Brittany coast on Europe's mainland and in southwest England.

populations of M. galloprovincialis occur in Brittany, France, and in Cornwall, England (Fig. 1). Thus, there are three northern limits of M. galloprovincialis in Europe. The North Sea, Irish Sea, and the eastern English Channel are composed almost exclusively of populations of M. edulis. A disjunct population of M. edulis occurs in a certain area of the Bay of Biscay in France (Fig. 1) where SST is generally cooler than the rest of the bay (Fig. 5A). We plotted these populations with respect to winter and summer SST, using February and August SST, respectively, to see how the distributions were related to the warmest and coldest environmental temperatures these populations experienced (Fig. 2). Populations of M. galloprovincialis in Europe never occur in areas where mean monthly SST falls below 9 °C in the winter, while populations of M. edulis occur in areas down to 6 °C (Fig. 2). In the summer, most populations of M. edulis occur where summer temperatures are below 20 °C, and no populations persist where summer SST is greater than 23 °C. Conversely, populations of M. galloprovincialis inhabit much warmer locations (Figure 2) including sites where summer SST regularly exceeds 23 °C. The warmest average summer temperature experienced by M. galloprovincialis in our dataset was 26.4 °C; however, the distribution of this species in northern Africa is not well-documented, so populations likely experience even warmer temperatures.

The Random Forest classification model, a correlative model, performed well, correctly classifying the out of bag samples (30% of data excluded when building the model) 97.14% of the time. Relative importance evaluation identified winter and spring variables (mean winter temperature, mean spring temperature, mean January temperature, mean March temperature, and mean April temperature) as the most important variables (ranked by mean decrease in model accuracy) for correctly classifying a site as Mytilus edulis or M. galloprovincialis. These results align nicely with a smaller-scale RF model based on M. galloprovincialis distribution in France and Spain, and explain historical changes in the distribution of M. galloprovincialis along the Normandy coast in France (Hilbish et al. 2012). Based on this correlative model and the known distribution of M. galloprovincialis with respect to SST, M. galloprovincialis is likely precluded from inhabiting areas where winter SST is

routinely below 9 °C, while *M. edulis* is more cold-adapted (Fig. 2). However, this correlative approach provides no explanation for the mechanism involved in this northern range limit of *M. galloprovincialis*. Previous work discerning the energetics of these two species (Fly and Hilbish 2013) provides evidence that the mechanism is not adult mussel energetics, as there are no significant differences at cold temperatures (Fig. 3).

Differences in energetics can, however, be ascertained as the mechanism for the differences in the species' southern limits. As stated previously, populations of *Mytilus edulis* do not inhabit areas where summer temperatures routinely exceed ~23 °C (Fig. 2). Fly and Hilbish (2013) determined the critical temperature at which SFG = 0 for *M. edulis* and *M. galloprovincialis*. While there were seasonal differences, the critical temperature for *M. edulis* was regularly around 23 °C (Fig. 3, Fly and Hilbish 2013), which corresponds with the environmental temperatures at the southern range edge of this species (Fig. 2, Fig. 4). The critical temperature for *M. galloprovincialis* was between 29–30 °C (Fig. 3, Fly and Hilbish 2013), and *M. galloprovincialis* inhabits regions in Europe with SST that does not exceed this threshold (Fig. 2, Fig. 4).

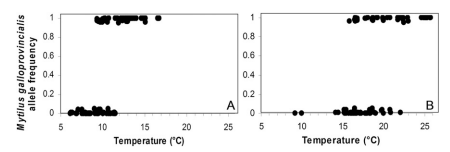


Figure 2. Distribution of European *Mytilus* populations with respect to winter and summer sea surface temperatures (SST). We plotted the populations shown in Figure 1 with respect to winter and summer SST, using **A**, February and **B**, August SST, respectively, to see how the distributions were related to environmental temperatures. **A**, Populations of *M. galloprovincialis* never occur in areas where mean monthly SST falls below 9 °C in the winter, while populations of *M. edulis* occur in areas down to 6 °C. **B**, In the summer, most populations of *M. edulis* occur where summer temperatures are below 20 °C, and no populations persist where summer SST is greater than 23 °C. Conversely, populations of *M. galloprovincialis* inhabit much warmer locations.

In the past decade, the 9 °C winter isotherm has been in the western part of the English Channel in the south (Fig. 5A). During the same period, the 20 °C summer isotherm has traversed the Bay of Biscay from just south of the mouth of the Gironde (45 °N) to NW Spain (Fig. 5B). This pattern is consistent with the exclusion of *Mytilus edulis* from the lower Bay of Biscay (Fig. 1). The EURO-CORDEX forecast predicts

both winter and summer warming (Fig. 5C-F). The result of this warming is the northward movement of the conditions for survival of M. edulis and M. galloprovincialis. The median of the EURO-CORDEX rcp4.5 forecasts predicts that summer maximum monthly SST of 20 °C will move northward to just south of the Brittany peninsula in France by 2050 (Fig. 5D), and move into the western English Channel and southern North Sea to 53°N by 2100 (Fig. 5F). This 20 °C isotherm is a proxy for the expected southern geographic limit of M. edulis, and predicts that the only region of the English Channel that will be habitable by M. edulis by 2100 will be along the Normandy coast of France (between Cherbourg and Calais). While a similar

permissible area appears on England's side of the English Channel (between Bournemouth and Dover), mussels are not readily found in this area due to a predominance of sandy beaches rather than hard substrates. The median of the rcp4.5 forecasts suggest that winter monthly SST higher than 9 °C will move eastward into the English Channel and northward into the Irish Sea by 2050 (Fig. 5C),

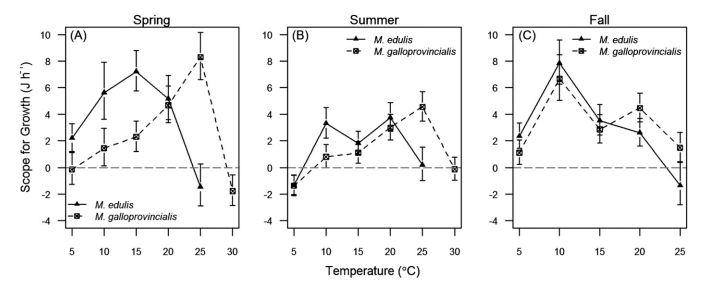


Figure 3. Physiological energetics of *Mytilus edulis* and *M. galloprovincialis*. Measurements of filtration rate, absorption efficiency and metabolic rate were used to derive an estimate of scope for growth (SFG) (sensu Widdows and Bayne 1971 the SFG measures the energy above maintenance available for growth and reproduction) of mussels at water temperatures ranging from 5 °C to 30 °C in **A,** spring, **B,** summer, and **C,** fall. No mussels survived at 30 °C in the fall. Estimates of the critical temperatures at which SFG = 0 (dashed line) were used to determine the geographic regions in Europe in which each species can potentially maintain a permanent population. This figure was adapted from Fly and Hilbish (2013).

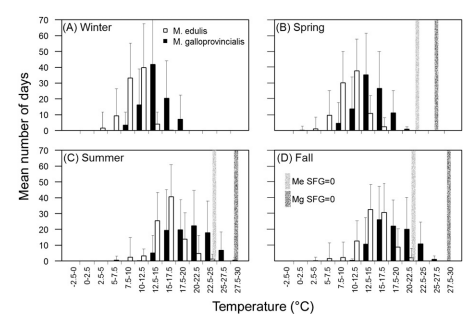


Figure 4. Average temperature profile of European *Mytilus edulis* (gray) and *M. galloprovincialis* (black) populations in each season. The transparent rectangles (gray for *M. edulis* and black for *M. galloprovincialis*) indicate the critical temperature at which SFG = 0 in spring, summer, and fall, as determined by Fly and Hilbish (2013). SFG data were not collected for winter ($\bf A$). In regions that exceed this critical temperature, populations of mussels are expected to be ephemeral or absent and, indeed, very few populations experience temperatures higher than that critical temperature. Summer ($\bf C$) and fall ($\bf D$) appear to be more physiologically-limiting than spring ($\bf B$). This figure was adapted from Fly and Hilbish (2013).

with only slightly further movement into the northern North Sea by 2100 (Fig. 5E). This prediction suggests that *M. galloprovincialis* will be much more prevalent in the British Isles and English Channel than it currently is.

DISCUSSION

Species distribution models are useful in understanding current and potential distributions of species; however, oftentimes one type of SDM is incapable of accurately predicting all of a species' range limits (e.g., north and south, east and west) (Woodin et al. 2013). This is because a given SDM applies a single mechanism or environmental profile across an entire species' range, when, it is often the case that different mechanisms or stressors control different portions of a species' geographic range or its distribution in different geographic regions (Sokolova et al. 2012, Woodin et al. 2013). This appears to be the case with Mytilus edulis and M. galloprovincialis. As suggested by mechanistic modeling, the southern limit of M. edulis is likely controlled by physiological constraints on the adult populations: specifically, populations cannot persist in areas where summer SST routinely

exceeds 20 °C, due to rapidly declining SFG above this temperature. It is likely that the southern limit of M. galloprovincialis is also controlled by energetic limitation; however, the southern limit of this species on the northwest African coast is not well-documented, so we cannot currently determine temperatures experienced at the very southern range edge of this species. Interestingly, while the southern limits of *Mytilus* spp. in Europe appear to be controlled by sublethal temperature effects of adult energetics, the southern limit of M. edulis on the east coast of the U.S. is controlled by lethal temperature effects on adult populations (Jones et al. 2010). This is likely because the east coast of the U.S. has a more continental climate (and, therefore, greater temperature extremes) than the west coast of Europe, which has a more coastal climate (Wethey 1985, Jenkins et al. 2008, Bailey 2009). Thus, mechanisms can differ within the same species in different regions.

The mechanism limiting the northern range of *Mytilus galloprovincialis* in Europe is still unknown; however, limi-

tations due to adult energetics are very unlikely. Blue mussels are physiologically quiescent during winter (Bayne 1976), and there are no major differences in the SFG of *M. edulis* and *M. galloprovincialis* at cold temperatures in other seasons that would explain the inability of *M. galloprovincialis* populations to persist where winter temperatures are below 9 °C (Fly and Hilbish 2013). This cold-water limitation has been noted previously for European *M. galloprovincialis* populations (Hilbish *et al.* 2012), as well as populations on the west coast of the U.S. (Hilbish *et al.* 2010). There are several hypotheses on the mechanism of this limitation that warrant further investigation, including tolerance to cold-shock and reproductive abilities.

The tolerance of species to cold temperatures depends on a variety of factors, including the thermal regimes and acclimation times experienced (Renault *et al.* 2004, Terblanche *et al.* 2011). Jansen *et al.* (2007) found that *Mytilus galloprovincialis* populations were much more sensitive to cold shock than *M. edulis* populations. This suggests that *M. galloprovincialis* might not have the same capability to recover from cumulative cold damage as *M. edulis*. Another possibility is that the reproductive capabilities of *M. galloprovincialis* are impeded at temperatures below 9 °C (Hilbish *et al.* 2010,

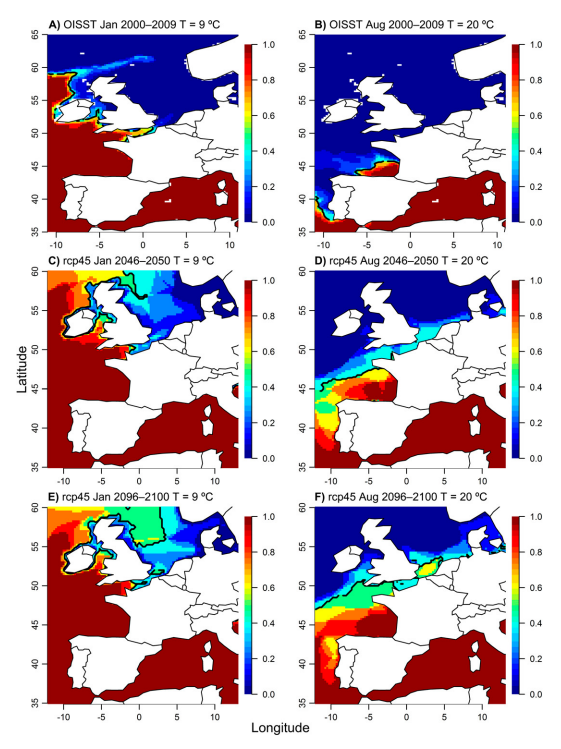


Figure 5. A, Fraction of winters 2002–2009 with temperatures above 9 °C in winter (February) and **B,** above 20 °C in summer (August), derived from NOAA Optimally Interpolated Sea Surface Temperature data. The contour line is the 50th percentile of the values throughout the figure. **C,** Fraction of EURO-CORDEX rcp4.5 models indicating temperatures above 9 °C in winter (February) and **D,** 20 °C in summer (August) in the period 2046–2050. **E,** Fraction of EURO-CORDEX rcp4.5 models indicating temperatures above 9 °C in winter (February) and **F,** 20 °C in summer (August) in the period 2096–2100. The temperature of 20 °C in summer is a proxy for the southern geographic limit of *Mytilus edulis* and the temperature of 9 °C is a proxy for the northern geographic limit of *M. galloprovincialis*. The contour lines for the forecasts (C–F) indicate the median of the model predictions. (Color shown in electronic version only).

Hilbish *et al.* 2012). Most populations of mussels spawn in the spring after having undergone gametogenesis over the winter (Bayne 1976). The fact that the RF model indicated winter or early spring as a key time for determining the distribution of these mussels suggests a hypothesis that some aspect of gametogenesis and/or spawning is temperature dependent and different between species.

Understanding the mechanism controlling this northern limit will be useful in making better predictions for the expansion of these populations as winter SSTs warm. The RF correlative model used monthly mean SST, while the mechanistic model considered distributions on a daily scale. Hilbish et al. (2012) ran a similar model using maximum entropy modeling (Maxent, http://www.cs.princeton.edu/~schapire/maxent/), that considered the number of days below temperature thresholds as variables. The two variables contributing most to mussel distribution were days below 9 °C and days below 10 °C (Hilbish et al. 2012). As hypotheses for mechanisms explaining this temperature distribution are examined, the temporal scale of the temperature distribution should be considered.

Warming of the ocean is not expected to occur uniformly (Xie et al. 2010), which suggests that the northern range limit of Mytilus galloprovincialis and the southern range limit of M. edulis will likely change at heterogeneous rates over the remainder of the 21st century. During the first half of the current century (present to 2050) the northern range limit of M. galloprovincialis is expected to shift substantially into the English Channel and into the Irish Sea and North Atlantic Ocean (Fig. 5C). However, during the second half of the century further range expansion is expected to be comparatively small (Fig. 5E). Conversely, the anticipated contraction of the southern range edge of M. edulis is expected to remain south of France's Brittany coast in the Bay of Biscay through 2050 (Fig. 5D), but shift dramatically north to 53°N by 2100, leaving only a small portion of French coastline, mainly the Normandy coast, suitable for *M. edulis* (Fig. 5F). France produces about 73,000 tons of mussels each year, 22% of which are produced in Normandy (Eurostat 2009). These potential future SST shifts indicate the bulk of the commercial mussel farms on the rest of the French coastline may no longer be able to cultivate *M. edulis* by the end of the century.

Mytilus edulis and M. galloprovincialis hybridize extensively, and the differing rates in their range expansions and contractions have implications for the hybrid zones formed between them. With a rapid northward expansion of M. galloprovincialis and minimal contraction of M. edulis by 2050, the hybrid zones between these two species could expand substantially. However, by 2099, many current regions of hybridization should be displaced by M. galloprovincialis and new hybrid zones may form in the Irish and North Atlantic Seas, regions currently inhabited primarily by M. edulis. This

brings to light the vulnerability of *M. edulis* to warming environmental conditions: those populations exposed to summer SSTs greater than 20 °C are highly vulnerable to being unable to persist.

Although we understand an important mechanism controlling the southern range edge of Mytilus edulis in Europe, we currently have still used 20 °C as our proxy for the physiological limit of this species. While the temperature at which SFG = 0 for M. edulis is \sim 23 °C, the southern range edge of this species maps more closely to a 20 °C SST isotherm, with only several populations persisting where summer temperatures range between 20-23 °C. It is likely that the populations living in this summer temperature range are physiologically stressed (Sokolova et al. 2012) and heavily dependent on other factors such as food availability. We should be able to develop an even stronger model to predict not only the range edge of this species with changing SST based on where SFG = 0, but to begin to quantify the productivity of these mussel beds based on the physiological rates measured (Fly and Hilbish 2013). These mechanistic models, as stated earlier, require much more input and background knowledge than correlative models. Thus, several issues must be resolved to produce a meaningful mechanistic model based on physiological productivity. A more detailed model will better take into account any differences between submerged and intertidal mussels. Mytilus edulis and M. galloprovincialis reduce their aerial rate of oxygen consumption to 4-17% that of their rate of aquatic oxygen consumption (Widdows et al. 1979), and incur only a small cost in terms of oxygen debt recovery (Widdows and Shick 1985). However, intertidal mussels can only feed while submerged, thus, affecting food availability and energy intake. A robust mechanistic model will take into account food availability and duration of exposure at low tide conditions.

The most outstanding question is how to incorporate food availability into projections of secondary production. Mussels consume food particles in the size range of 2–20 µm (Bayne 1976), but algal biomass within this size range is only, at best, weakly correlated with chlorophyll a or other pigments that can be assessed with remote sensing (Alpine and Cloern 1985, Han and Furuya 2000, Arin et al. 2002). Additionally remote sensing products are often unreliable in near-shore coastal environments where materials of terrestrial origin and reflection off of the ocean bottom interfere with sensor readings (Hellweger et al. 2004, Moses et al. 2009). The result is that it is very difficult to predict present day secondary production of mussels (and many other marine species) on a regional scale. It will be even more difficult to forecast changes in primary production, specifically of the nanoplankton as a consequence of climate change and to incorporate these changes into models of secondary production of mussels and other coastal species (Sommer and

Lengfellner 2008). Until meaningful projections of changes in coastal primary production become available the most productive approach may be to forecast changes in local and regional potential for secondary production and provide mechanistic models that will allow the incorporation of local primary production to assess the management of coastal species with respect to climate change.

ACKNOWLEDGMENTS

This study was supported by grants from NSF (OCE1039513, OCE1129401), and NASA (NNX07AF20G, NNX11AP77G). We acknowledge the World Climate Research Programme's Working Group on Regional Climate, and the Working Group on Coupled Modelling, former coordinating body of CORDEX and responsible panel for CMIP5. We also thank the climate modelling groups (listed in Appendix B of this paper) for producing and making available their model output. We also acknowledge the Earth System Grid Federation infrastructure, an international effort led by the U.S. Department of Energy's Program for Climate Model Diagnosis and Intercomparison, the European Network for Earth System Modelling and other partners in the Global Organisation for Earth System Science Portals (GO-ESSP). P. M. Brannock, A. O. Hollis, and S. J. Jones assisted with field and genetic work. We are grateful to P. B. Marko and B. A. Seibel, the American Malacological Society, and the University of the Azores for their organization of the World Congress of Malacology 2013 conference and the symposium on "Climate Change and Molluscan Ecophysiology." This is publication number 73 in Ecological Forecasting from the University of South Carolina.

LITERATURE CITED

- Akima, H., A. Gebhardt, and T. Petzoldt. 2013. Akima: Interpolation of irregularly spaced data. R. package version 0.5-10. Internet resource: http://CRAN.R-project.org/package=akima.
- Alpine, A. E. and J. E. Cloern. 1985. Differences in *in vivo* fluorescence yield between three phytoplankton size classes. *Journal of Plankton Research* 7: 381–390.
- Arin, L., X. A. G. Moran, and M. Estrada. 2002. Phytoplankton size distribution and growth rates in the Alboran Sea (SW Mediterranean): Short term variability related to mesoscale hydrodynamics. *Journal of Plankton Research* 24: 1019–1033.
- Bailey, R. G. 2009. Ecoregions of the United States. In R. G. Bailey, ed., Ecosystem Geography: From Ecoregions to Sites. 2nd edition. Springer-Verlag. New York, New York.
- Banni, M., A. Negri, F. Mignone, H. Boussetta, A. Viarengo, and F. Dondero. 2011. Gene expression rhythms in the mussel *Mytilus galloprovincialis* (Lam.) across an annual cycle. *PLoS ONE* **6**: e18904.

- Bayne, B. L. 1976. *Marine Mussels: Their Ecology and Physiology*. Cambridge University Press, Cambridge, United Kingdom.
- Bierne, N., P. Borsa, C. Daguin, D. Jollivet, F. Viard, F. Bonhomme, and P. David. 2003. Introgression patterns in the mosaic hybrid zone between *Mytilus edulis* and *M. galloprovincialis*. *Molecular Ecology* **12**: 447–461.
- Breiman, L. 2001. Random forests. Machine Learning 45: 5-32.
- Buckley, L. B., M. C. Urban, M. J. Angilletta, L. G. Crozier, L. J. Rissler, and M. W. Sears. 2010. Can mechanism inform species' distribution models? *Ecology Letters* 13: 1041–1054.
- Burrows, M, T., D. S. Schoeman, L. B. Buckley, P. Moore, E. S. Poloczanska, K. M. Brander, C. Brown, J. F. Bruno, C. M. Duarte, B. S. Halpern, J. Holding, C. V. Kappel, W. Kiessling, M. I. O'Connor, J. M. Pandolfi, C. Parmesan, F. B. Schwing, W. J. Sydeman, and A. J. Richardson. 2011. The pace of shifting climate in marine and terrestrial ecosystems. *Science* 334: 652–655.
- Coustau, C., F. Renaud, and B. Delay. 1991. Genetic characterization of the hybridization between *Mytilus edulis* and *M. galloprovincialis* on the Atlantic coast of France. *Marine Biology* 111: 87–93.
- Cutler, D. R., T. C. Edwards, K. H. Beard, A. Cutler, K. T. Hess, J. Gibson, and J. J. Lawler. 2007. Random forests for classification in ecology. *Ecology* 88: 2783–2792.
- Daguin, C., F. Bonhomme, and P. Borsa. 2001. The zone of sympatry and hybridization of *Mytilus edulis* and *M. galloprovincialis*, as described by intron length polymorphism at locus *mac-1*. *Heredity* **86**: 342–354.
- Edwards, M. and A. J. Richardson. 2004. Impact of climate change on marine pelagic phenology and trophic mismatch. *Nature* 430: 881–884.
- Elith, J. and J. R. Leathwick. 2009. Species distribution models: Ecological explanation and prediction across space and time. *Annual Review of Ecology, Evolution, and Systematics* **40**: 677–697.
- Eurostat, 2009. Eurostat statistics in focus. Agriculture and Fisheries, 83/2009.
- FAO. 2012. Cultured Aquatic Species Information Programme. *Mytilus edulis*. Text by Goulletquer, P. *In*: FAO Fisheries and Aquaculture Department [online]. Rome, Italy. http://www.fao.org/fishery/culturedspecies/Mytilus_edulis/ 6 February 2012.
- Fly, E. K. and T. J. Hilbish. 2013. Physiological energetics and biogeographic range limits of three congeneric mussel species. *Oecologia* 172: 35–46.
- Gardner, J. P. A. and D. O. F. Skibinski. 1991. Biological and physical factors influencing genotype-dependent mortality in hybrid mussel populations. *Marine Ecology Progress Series* 71: 235–243.
- Gilg, M. R. and T. J. Hilbish. 2003. The geography of marine larval dispersal: Coupling genetics with fine-scale physical oceanography. *Ecology* **84**: 2989–2998.
- Gosling, E., S. Doherty, and N. Howley. 2008. Genetic characterization of hybrid mussel (*Mytilus*) populations on Irish coasts. *Journal of the Marine Biological Association of the UK* **88**: 341–346.
- Guisan, A. and W. Thuiller. 2005. Predicting species distribution: Offering more than simple habitat models. *Ecology Letters* 8: 993–1009.

- Han, M.-S. and K. Furuya. 2000. Size and species-specific primary productivity and community structure of phytoplankton in Tokyo Bay. *Journal of Plankton Research* 22: 1221–1235.
- Hellweger, F. L., P. Schlosser, U. Lall, and J. K. Weissel. 2004. Use of satellite imagery for water quality studies in New York Harbor. *Estuarine, Coastal and Shelf Science* **61**: 437–448.
- Helmuth, B., J. G. Kingsolver, and E. Carrington. 2005. Biophysics, physiological ecology, and climate change: Does mechanism matter? *Annual Review of Physiology* **67**: 177–201.
- Hilbish, T. J., P. M. Brannock, K. R. Jones, A. B. Smith, B. N. Bullock, and D. S. Wethey. 2010. Historical changes in the distributions of invasive and endemic marine invertebrates are contrary to global warming predictions: The effects of decadal climate oscillations. *Journal of Biogeography* 37: 423–431.
- Hilbish, T. J., E. W. Carson, J. R. Plante, L. A. Weaver, and M. R. Gilg. 2002. Distribution of *Mytilus edulis, M. galloprovincialis* and their hybrids in open-coast populations of mussel in southwestern England. *Marine Biology* 140: 137–142.
- Hilbish, T. J., F. P. Lima, P. M. Brannock, E. K. Fly, R. L. Rognstad, and D. S. Wethey 2012. Change and stasis in marine hybrid zones in response to climate warming. *Journal of Biogeography* 39: 676–687.
- Jacob, D., J. Petersen, B. Eggert, A. Alias, O. B. Christensen, L. M. Bouwer, A. Braun, A. Colette, M. Déqué, G. Georgievski, E. Georgopoulou, A. Gobiet, L. Menut, G. Nikulin, A. Haensler, N. Hempelmann, C. Jones, K. Keuler, S. Kovats, N. Kröner, S. Kotlarski, A. Kriegsmann, E. Martin, E. van Meijgaard, C. Moseley, S. Pfeifer, S. Preuschmann, C. Radermacher, K. Radtke, D. Rechid, M. Rounsevell, P. Samuelsson, S. Somot, J.-F. Soussana, C. Teichmann, R. Valentini, R. Vautard, B. Weber, and P. Yiou. 2014. EURO-CORDEX: New high-resolution climate change projections for European impact research. Regional Environmental Change 14: 563-578.
- Jansen, J. M., S. W. Bonga, and H. Hummel. 2007. Differential coldshock resistance among acclimated European mussel populations. *Marine and Freshwater Behaviour and Physiology* 40: 233–245.
- Jenkins, S. R., P. Moore, M. T. Burrows, D. J. Garbary, S. J. Hawkins, A. Ingólffson, K. P. Sebens, P. V. R. Snelgrove, D. S. Wethey, and S. A. Woodin. 2008. Comparative ecology of North Atlantic shores: Do differences in players matter for process? *Ecology* 89: S3–S23.
- Jones, S. J., F. P. Lima, and D. S. Wethey. 2010. Rising environmental temperatures and biogeography: Poleward range contraction of the blue mussel, *Mytilus edulis* L., in the western Atlantic. *Journal of Biogeography* 37: 2243–2259.
- Jones, S. J., A. J. Southward, and D. S. Wethey. 2012. Climate change and historical biogeography of the barnacle *Semibalanus balanoides*. *Global Ecology and Biogeography* **21**: 716–724.
- Kearney, M. and W. Porter. 2009. Mechanistic niche modeling: Combining physiological and spatial data to predict species' ranges. *Ecology Letters* **12**: 334–350.
- Kijewski, T., B. Śmietanka, M. Zbawkicka, E. Gosling, H. Hummel, and R. Wenne. 2011. Distribution of *Mytilus* taxa in European coastal areas as inferred from molecular markers. *Journal of Sea Research* **65**: 224–234.
- Lawler, J. J., D. White, R. P. Neilson, and A. R. Blaustein. 2006. Predicting climate-induced range shifts: Model differences and model reliability. *Global Change Biology* **12**: 1568–1584.

- Liaw, A. and M. Wiener. 2002. Classification and regression by randomForest. *R News* 2/3: 18–22.
- Lionetto, M. G., R. Caricato, M. E. Giordano, M. F. Pascariello, L. Marinosci, and T. Schettino. 2003. Integrated use of biomarkers (acetylcholinesterase and antioxidant enzymes activities) in *Mytilus galloprovincialis* and *Mullus barbatus* in an Italian coastal marine area. *Marine Pollution Bulletin* **46**: 324–330.
- McDonald, J. H., R. Seed, and R. K. Koehn. 1991. Allozymes and morphometric characters of three species of *Mytilus* in the Northern and Southern Hemispheres. *Marine Biology* 111: 323–333.
- Moses, W. J., A. A. Gitelson, S. Berdnikov, and V. Povazhnyy. 2009. Estimation of chlorophyll-*a* concentration in case II waters using MODIS and MERIS data successes and challenges. *Environmental Research Letters* **4**: 045005, doi:10.1088/1748-9326/4/4/045005.
- Peterson, A. T., J. Soberón, R. G. Pearson, R. P. Anderson, E. Martínez-Meyer, M. Nakamura, and M. B. Araújo. 2011. Ecological Niches and Geographic Distributions. Princeton University Press, Princeton, New Jersey, U.S.A.
- Philippart, C. J. M., H. M. van Aken, J. J. Beukema, O. G. Bos, G. C. Cadée, and R. Dekker. 2003. Climate-related changes in recruitment of the bivalve *Macoma balthica*. *Limnology and Oceanography* 48: 2171–2185.
- Pisanelli, B., M. Benedetti, D. Fattorini, and F. Regoli. 2009. Seasonal and inter-annual variability of DNA integrity in mussels *Mytilus galloprovincialis*: A possible role for natural fluctuations of trace metal concentrations and oxidative biomarkers. *Chemosphere* 77: 1551–1557.
- R Development Core Team. 2010. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
- Rawson, P. D. and T. J. Hilbish. 1998. Asymmetric introgression of mitochondrial DNA among European populations of blue mussels (*Mytilus* spp.). *Evolution* **52**: 100–108.
- Renault, D., O. Nedved, F. Hervant, and P. Vernon. 2004. The importance of fluctuating thermal regimes for repairing chill injuries in the tropical beetle *Alphitobius diaperinus* (Coleoptera: Tenebrionidae) during exposure to low temperature. *Physiological Entomology* **29**: 139–145.
- Reynolds, R. W., T. M. Smith, C. Liu, D. B. Chelton, K. S. Casey, and M. G. Schlax. 2007. Daily high-resolution-blended analysis for sea surface temperature. *Journal of Climate* 20: 5473–5496.
- Roberston, M. P., C. I. Peter, M. H. Villet, and B. S. Ripley. 2003. Comparing models for predicting species' potential distributions: a case study using correlative and mechanistic predictive modeling techniques. *Ecological Modelling* **164**: 153–167.
- Sará, G., M. Kearney, and B. Helmuth. 2011. Combining heattransfer and energy budget models to predict thermal stress in Mediterranean intertidal mussels. *Chemistry and Ecology* 27: 135–145.
- Seed, R.1992. Systematics evolution and distribution of mussels belonging to the genus *Mytilus*: An overview. *American Malacological Bulletin* **9**: 123–137.
- Skibinski, D. O. F., J. A. Beardmore, and T. F. Cross. 1983. Aspects of the population genetics of *Mytilus* (Mytilidae; Mollusca) in the British Isles. *Biological Journal of the Linnean Society* **19**: 137–183.

- Sokolova, I. M., M. Frederich, R. Bagwe, G. Lannig, and A. A. Sukhotin. 2012. Energy homeostasis as an integrative tool for assessing limits of environmental stress tolerance in aquatic invertebrates. *Marine Environmental Research* 79: 1–15.
- Sommer, U. and K. Lengfellner. 2008. Climate change and the timing, magnitude, and composition of the phytoplankton spring bloom. *Global Change Biology* **14**: 1199–1208.
- Suchanek, T. H. 1992. Extreme biodiversity in the marine environment: Mussel bed communities of Mytilus californianus. Northwest Environmental Journal 8: 150–152.
- Terblanche, J. S., A. A. Hoffmann, K. A. Mitchell, L. Rako, P. C. le Roux, and S. L. Chown. 2011. Ecologically relevant measures of tolerance to potentially lethal temperatures. *Journal of Experimental Biology* **214**: 3713–3725.
- Wethey, D. S. 1985. Catastrophe, extinction and species diversity: A rocky intertidal example. *Ecology* **66**: 445–456.
- Widdows, J. and B. L. Bayne. 1971. Temperature acclimation of Mytilus edulis with reference to its energy budget. Journal of the Marine Biological Association of the UK 51: 827–843.
- Widdows, J., P. Fieth, and C. M. Worrall. 1979. Relationships between seston, available food and feeding activity in the common mussel *Mytilus edulis*. *Marine Biology* **50**: 195–207.
- Widdows, J. and J. M. Shick. 1985. Physiological responses of Mytilus edulis and Cardium edule to aerial exposure. Marine Biology 85: 217–232.
- Woodin, S. A., T. J. Hilbish, B. Helmuth, S. J. Jones, and D. S. Wethey. 2013. Climate change, species distribution models, and physiological performance metrics: Predicting when biogeographic models are likely to fail. *Ecology and Evolution* 3: 3334–3346.
- Xie, S.-P., C. Deser, G. A. Vecchi, J. Ma, H. Teng, and A. T. Wittenberg, 2010. Global warming pattern formation: Sea surface temperature and rainfall. *Journal of Climate* 23: 966–986.

Submitted: 10 December 2013; accepted 31 August, 2014; final revisions received: 15 January 2015

Appendix A Data used for Europe-wide Random Forest model using distributions of "pure" populations of *Mytilus galloprovincialis* and *M. edulis* based on published accounts or genetic analyses done for this study where *M. galloprovincialis* allele frequencies were greater than 95% or less than 5%, respectively. The sample sites are listed by latitude from North to South.

Year sampled	Site	Latitude	Longitude	Frequency of Mg allele	Source
2003	White Sea, Russia	66.00	39.37	0.00	Kijewski et al. 2011
2003	Reykjavik, Iceland	64.17	-22.02	0.00	Kijewski et al. 2011
2008	Tayport, Scotland	56.44	-2.94	0.01	current study
2008	Cramond, Scotland	55.98	-3.30	0.01	current study
2009	Fairlie, Scotland	55.76	-4.86	0.04	current study
2009	Millport, Scotland	55.75	-4.93	0.01	current study
2009	Rockcliffe, England	54.85	-3.79	0.00	current study
2009	Seascale, England	54.40	-3.49	0.00	current study
2009	Ravenglass, England	54.36	-3.41	0.00	current study
2008	Filey Brigg South, England	54.22	-0.27	0.04	current study
2005	Carlingford, Ireland	54.04	-6.18	0.00	Gosling et al. 2008
2005	Dunany, Ireland	53.86	-6.24	0.00	Gosling et al. 2008
2009	Thurstaston, England	53.34	-3.15	0.00	current study
2005	Dublin, Ireland	53.30	-6.15	0.00	Gosling et al. 2008
2009	Conwy, Wales	53.29	-3.83	0.00	current study
2008	Heacham, England	52.91	0.47	0.00	current study
2008	Lowestoft, England	52.45	1.74	0.01	current study
2008	Llangranog, Wales	52.16	-4.47	0.01	current study
2008	Harwich, England	51.93	1.27	0.00	current study
2008	Solva, Wales	51.87	-5.20	0.00	current study
2008	Milford Haven, Wales	51.72	-5.10	0.02	current study
2008	Dales, Wales	51.71	-5.17	0.00	current study
2008	Burry Port, Wales	51.68	-4.25	0.00	current study
2008	South Woodham Ferrers, England	51.63	0.61	0.00	current study
2008	Port Talbot, Wales	51.59	-3.81	0.00	current study
2008	Mumbles, Wales	51.57	-3.99	0.00	current study
2008	Porthcawl, Wales	51.50	-3.74	0.00	current study
2008	Atlantic College, England	51.40	-3.53	0.00	current study
2008	Swale, England	51.35	0.88	0.00	current study
2008	Minehead, England	51.21	-3.46	0.00	current study
2009	Exmouth, England	50.61	-3.41	0.00	current study
2009	Teignmouth, England	50.54	-3.49	0.00	current study
2009	Maidencombe, England	50.51	-3.51	0.00	current study
2009	Torbay, England	50.45	-3.54	0.00	current study
2009	Dartmouth, England	50.34	-3.57	0.00	current study
2009	Torcross, England	50.26	-3.65	0.00	current study
2003	Vlissengen, Netherlands	51.44	3.56	0.05	Kijewski <i>et al.</i> 2011
2003	Oostende, Belgium	51.24	2.92	0.00	Kijewski <i>et al.</i> 2011
2005	Dunkirk, France	51.24	2.38	0.00	Hilbish <i>et al.</i> 2012
	Dieppe, France			0.00	Hilbish et al. 2012
2005		49.94	1.09		
2005	Saint-Vaast-la-Hougue, France	49.58	-1.26	0.00	Hilbish et al. 2012
2003	Seine, France	49.42	0.08	0.00	Kijewski <i>et al.</i> 2011
2007	Merville-Franceville, France	49.29	-0.20	0.00	Hilbish et al. 2012
2006	Billiers, France	47.52	-2.48	0.04	Hilbish et al. 2012
2006	Barbatre, France	46.89	-2.15	0.01	Hilbish et al. 2012
2005	Saint-Gilles-Croix-de-Vie, France	46.69	-1.95	0.04	Hilbish et al. 2012
2006	La Tranche-sur-Mer, France	46.34	-1.46	0.03	Hilbish et al. 2012
2006	Vieux Châtelaillon, France	46.06	-1.09	0.00	Hilbish et al. 2012
2006	Montalivet-les-Bains, France	45.38	-1.16	0.01	Hilbish et al. 2012
2006	Mimizan-Plage, France	44.21	-1.29	0.01	Hilbish et al. 2012
2006	Pointe de Grave, France	45.57	-1.06	0.04	Hilbish et al. 2012
2006	Pornic, France	47.11	-2.11	0.05	Hilbish et al. 2012

Appendix A. (Continued)

Year sampled	Site	Latitude	Longitude	Frequency of Mg allele	Source
2006	Saint-Brevin-les-Pins, France	47.27	-2.17	0.03	Hilbish et al. 2012
2006	Nacqueville, France	49.68	-1.72	0.01	Hilbish et al. 2012
2006	Pointe du Moulard, France	49.64	-1.23	0.00	Hilbish et al. 2012
2006	Ravenoville, France	49.47	-1.24	0.00	Hilbish et al. 2012
2006	Sainte-Honorine-des-Pertes, France	49.36	-0.80	0.00	Hilbish et al. 2012
2006	Luc-sur-Mer, France	49.32	0.47	0.00	Hilbish et al. 2012
2006	Saint-Jouin-Bruneval, France	49.65	0.15	0.00	Hilbish et al. 2012
2006	Étretat, France	49.71	0.20	0.00	Hilbish et al. 2012
2006	Saint-Pierre-en-Port, France	49.81	0.49	0.01	Hilbish et al. 2012
2006	Saint-Valéry-en-Caux, France	49.87	0.71	0.00	Hilbish et al. 2012
2006	Pourville, France	49.92	1.03	0.00	Hilbish et al. 2012
2009	Port Quin, England	50.59	-4.87	0.99	current study
2009	Rock, England	50.54	-4.92	0.98	current study
2009	Porthcothan, England	50.51	-5.03	0.97	current study
2009	Newquay, England	50.42	-5.07	0.96	current study
2009	Trevaunance Cove, England	50.32	-5.20	0.98	current study
2009	Portreath, England	50.26	-5.30	0.98	current study
2009	Saint Ives, England	50.22	-5.49	0.96	current study
2005	Île Callot, France	48.68	-3.92	0.98	Hilbish et al. 2011
2005	Roscoff, France	48.73	-3.99	1.00	Hilbish et al. 2012
2005	Île Grande, France	48.81	-3.57	0.99	Hilbish et al. 2013
2007	Locquirec, France	48.69	-3.64	0.99	Hilbish et al. 2012
2007	Brignogan-Plage, France	48.67	-4.32	0.96	Hilbish et al. 2012
2006	Saint-Jean-de-Luz, France	43.39	-1.67	1.00	Hilbish et al. 2012
2005	Viveiro, Spain	43.72	-7.62	0.97	Hilbish et al. 2012
2005	Ortigueira, Spain	43.71	-7.85	0.99	Hilbish et al. 2012
2005	Foz, Spain	43.57	-7.25	0.98	Hilbish et al. 2012
2006	Laredo, Spain	43.41	-3.42	1.00	Hilbish et al. 2012
2005	Cabanas, Spain	43.41	-8.17	0.99	Hilbish et al. 2012
2006	San Vincente de la Barquera, Spain	43.39	-4.37	0.98	Hilbish et al. 2012
2006	Hondarribia, Spain	43.37	-1.79	0.95	Hilbish et al. 2012
2005	A Coruña, Spain	43.37	-8.38	0.98	Hilbish et al. 2012
2006	Deba, Spain	43.30	-2.35	0.98	Hilbish et al. 2012
2005	Laxe, Spain	43.22	-9.00	0.97	Hilbish et al. 2012
2005	Esteiro, Spain	42.79	-8.98	0.99	Hilbish et al. 2012
2005	Marin, Spain	42.40	-8.69	0.99	Hilbish et al. 2012
2005	San Xenxo, Spain	42.38	-8.85	0.99	Hilbish et al. 2012
2005	Bueu, Spain	42.23	-8.79	0.99	Hilbish et al. 2012
2005	Viana do Castelo, Portugal	41.70	-8.86	1.00	Hilbish et al. 2012
2005	Ílhavo, Portugal	40.62	-8.75	1.00	Hilbish et al. 2012
2005	São Martinho do Porto, Portugal	39.52	-9.15	1.00	Hilbish et al. 2012
2005	Tróia, Portugal	38.49	-8.90	1.00	Hilbish et al. 2012
2006	Sines, Portugal	37.91	-8.80	1.00	Hilbish et al. 2012
2005	Albufeira, Portugal	37.08	-8.23	1.00	Hilbish et al. 2012
2006	Burgau, Portugal	37.07	-8.77	1.00	Hilbish et al. 2012
2003	Cádiz, Spain	36.55	-6.37	0.96	Kijewski et al. 2011
2006	Ancona, Italy	43.56	13.59	1.00	Pisanelli et al. 2009
2006	Cap d'Agde, France	43.27	3.52	1.00	Wethey, unpubl. da
2003	Banyuls-sur-Mer, France	42.48	3.13	1.00	Kijewski et al. 2011
2003	Barcelona, Spain	41.37	2.20	1.00	Kijewski et al. 2011
2000	Carovigno, Italy	40.72	17.80	1.00	Lionetto et al. 2003
2000	Brindisi, Italy	40.65	17.98	1.00	Lionetto et al. 2003
2000	Torchiarolo, Italy	40.54	18.08	1.00	Lionetto et al. 2003
2006	Peñíscola, Spain	40.36	0.40	1.00	Wethey, unpubl. dat

Appendix A. (Continued)

Year sampled	Site	Latitude	Longitude	Frequency of Mg allele	Source
2000	Lecce, Italy	40.34	18.37	1.00	Lionetto et al. 2003
2000	Otranto, Italy	40.15	18.49	1.00	Lionetto et al. 2003
2000	Santa Maria al Bagno, Italy	40.13	17.99	1.00	Lionetto et al. 2003
2006	Oropesa del Mar, Spain	40.08	0.14	1.00	Wethey, unpubl. data
2000	Gallipoli, Italy	40.05	17.97	1.00	Lionetto et al. 2003
2000	Castro, Italy	40.00	18.43	1.00	Lionetto et al. 2003
2003	Oristano, Sardinia	39.86	8.55	1.00	Kijewski et al. 2011
2008	Bizerte, Tunisia	37.27	9.89	1.00	Banni et al. 2011
2006	Torrox, Spain	36.73	-3.96	1.00	Wethey, unpubl. data
2006	Estepona, Spain	36.41	-5.17	1.00	Wethey, unpubl. data

Appendix B. Combinations of global and regional circulation models used in ensemble predictions of ocean climate change. C4I, Community Climate Change Consortium for Ireland, CNRM, Météo France, DMI, Danish Meteorological Institute, ETHZ, Swiss Institute of Technology Zurich, GKSS, Helmholz Center Geesthacht Institute for Coastal Research, ICTP, KNMI, Royal Netherlands Meteorological Institute, HC, UK Met Office Hadley Centre, MPI, Max Planck Institute, OURANOS, Consortium on Regional Climatology and Adaptation to Climate Change, SMHI, Swedish Meteorological and Hydrological Institute, UCLM, Universidad Castilla, La Mancha, VMGO, Voeikov Main Geophysical Observatory.

Modeling Group	Global Model	Regional Model
C4I	ECHAM5	RCA3
CNRM	ARPEGE	Aladin4.5
CNRM	ARPEGE	Aladin5.1
DMI	ARPEGE	HIRHAM5
DMI	ECHAM5	HIRHAM5
DMI	BCM	HIRHAM5
ETHZ	HadCM3Q0	CLM
GKSS	IPSL	CLM
KNMI	ECHAM5-r3	RACMO2
HC	HadCM3Q0	HadRM3Q0
HC	HadCM3Q3	HadRM3Q3
HC	HadCM3Q16	HadRM3Q16
MPI	ECHAM5	M-REMO
OURANOS	CGCM3	CRCM
SMI	BCM	RCA
SMI	ECHAM5-r3	RCA
SMI	HadCM3Q3	RCA
UCLM	HadCM3Q0	PROMES
VMGO	HadCM3Q0	RRCM