INTRODUCTION

The highest mortality rates in most marine organisms occur during their early life stages (McGurk 1986), so factors affecting the placement of eggs and the maturation of larval stages are critical aspects of a species’ life history strategy. It is well established that in marine organisms, the adaptation to regional environments reflects adult responses for when and where to spawn that tend to optimize the offspring survival (Iles & Sinclair 1982). However, the cost associated with reproductive activities, such as migration and gonad maturity, are also an important factor constraining the spawning locations of a species.

Many marine species undertake long migrations to reach their spawning grounds. Annual spawning migrations in temperate marine species are complex processes that result from the interactions among physiology, behavior, adult life history strategies, and larval ecology (Cury 1994, Corten 2001, Jørgensen et al. 2008). In contrast, other species have fairly restric-
ted dispersal abilities and complete their life cycle in relatively closed spatial areas. It is possible that within similar geographical ranges, species with different life history traits can reproduce and thrive, but also show adaptations to local spawning habitats at smaller spatial scales.

The cues and clues that fish respond to in their decisions regarding when and where to spawn can be geographically and environmentally based (Bai-ley et al. 2004, Ciannelli et al. 2007). Some species may adapt to spawn in geographic areas where the environment tends to be more stable, and where habitat characteristics are optimal for the survival of later larval and juvenile stages. Other species may adapt toward a more opportunistic strategy and spawn within specific environmental ranges of a larger geographic area. Depending on the physical tolerances of the species, the 2 strategies can show any degree of mixing. At a regional spatial scale (e.g. around the Balearic Islands), the prevalence of one or the other spawning strategy can be correlated with specific life history traits. The important question then is how species’ life history traits and spawning strategies are linked. Providing an answer to this question would allow us to generate hypotheses on how spawning strategies are regulated across a wide range of species, and thus to determine their importance in fisheries management decisions.

In this study, we were particularly interested in contrasting spawning strategies of closely related and spatially co-occurring species. We compared the degree to which spawning locations are either environmentally or geographically fixed in 3 species of tuna with contrasting life history strategies, but all spawning within the Balearic Islands region in the NW Mediterranean Sea. We focused our analyses on a larval tuna data set collected over 5 yr (2001 to 2005). The larva data cover a wide geographical scale (180 × 220 nautical miles) in the Balearic area of the Mediterranean Sea, where large spawning aggregations of tuna and other pelagic predator species are observed during spring and summer. The 3 tuna species that we investigated are Atlantic bluefin tuna *Thunnus thynus*, Mediterranean albacore *T. alalunga*, and bullet tuna *Auxis rochei*. Part of the population of the large migratory Atlantic bluefin tuna comes from the Atlantic and crosses the Strait of Gibraltar to reach the Balearic Islands to spawn (Block et al. 2005, Rooker et al. 2008). This species is a large, oceanic temperate tuna. The Mediterranean albacore is a smaller tropical tuna that ranges widely in the Mediterranean Sea through its life cycle, but reproduces in large aggregations near the Balearic Islands (Arrizabalaga et al. 2004). Adult bullet tuna, also a small tropical species, spawn and reside in more coastal locations (Sabatés & Recasens 2001).

**MATERIALS AND METHODS**

The study area is the Balearic archipelago in the western Mediterranean basin. This area is characterized by the advection of new Atlantic water from the south that converges with northern resident Atlantic water, forming oceanographic structures such as fronts (Fig. 1). In summer, larval assemblages around the Balearic Islands show high proportions of the 3 tuna species examined in this study. The spatial distribution of spawning for these 3 selected tuna species was inferred from their early-larvae distributions. During June and July 2001 to 2005, tuna fish larvae were collected using a 333 µm mesh fitted to 60 cm mouth opening Bongo nets that were equipped with flowmeters. Oblique tows were performed down to 70 m in the open sea or down to 5 m above the sea floor in shallower areas. Samples from the 333 µm net were fixed with 4% formalin buffered with borax, and the larvae of the tuna species were further identified in the laboratory. Standard length (SL, mm) of all tuna larvae captured was measured without correction for shrinkage, and those identified as yolk sac and pre-flexion developmental stages (<4.5 mm) were merged into a class denominated ‘stage-1.’ Most studies using tuna larval data described spawning distribution for all developmental stages pooled together (Alemany et al. 2010, Muhling et al. 2010). The duration of the yolk-sac developmental stage in these larvae is very short (up to 1 d, Margulies et al. 2007). Since dispersal mechanisms are important processes affecting the distribution of these early larval stages, we selected only the yolk-sac and the pre-flexion life stages. Around 200 stations located in a regular grid of 10 × 10 nautical miles were sampled every year (Table 1). This spatial resolution was chosen to take into account the patchy distribution of tuna species in the sampling (Alemany et al. 2010, Satoh 2010).

Among other hydrographic measurements, water temperature and salinity were recorded *in situ* at all stations using a CTD down to 350 m and at some deep stations down to 650 m (for more information on the variables measured, see Alemany et al. 2010). The geostrophic velocity (measured in cm s⁻¹) was estimated by differentiating the interpolated dynamic height field calculated from CTD measurements as described by Torres et al. (2011).
We developed 2 nonparametric regression statistical models (generalized additive models, GAMs; Wood 2006) to assess (1) spatially varying patterns of spawning among the 3 species and (2) the importance of environmental variability in driving the locations for spawning. In all cases, the dependent variable was the number of stage-1 larvae of each species counted in the sample (Stage1). The covariates of the first GAM included latitude (lat) and longitude (lon) of the sampling stations to incorporate into the model the geographic effect, day of the year (cd) to address the timing of spawning, and hour of sampling (hour) to account for differences in larval catchability along the day cycle. GAMs based on these variables were used to describe geographical patterns in spawning alone, hereafter referred to as the ‘geographical model.’ The formulation of the geographical model was:

\[
\text{Stage1}_{\text{species}} = a_y + \text{offset} \left( \log(\text{vol}) \right) + \text{factor(year)} + s_1(\text{lat, lon}) + s_2(\text{cd}) + s_3(\text{hour}) + \varepsilon
\]  

(1)

where \( s \) is either a 2-dimensional \( s_1 \) or a unidimensional \( s_2, s_3 \) smoothing function, typically a thin plate regression spline (Wood 2006), \( a_y \) is a year-dependent model intercept, and \( \varepsilon \) is the error. Due to the highly patchy nature of the abundances, we used an over-dispersed Poisson distribution to model larval data and a natural-log link. Volume of water filtered (vol, m\(^3\)) was included as an offset (after natural log transformation), to account for the effort expended in catching the sample. Sampling years were included as factors in the model to account for interannual variations of spawning output (related to spawning stock size).

Table 1. Auxis rochei, Thunnus alalunga, and T. thynnus. Number of stations where larvae of the 3 tuna species were found and total number of stations analyzed each year.

<table>
<thead>
<tr>
<th>Year</th>
<th>A. rochei</th>
<th>T. alalunga</th>
<th>T. thynnus</th>
<th>No. stations analyzed</th>
</tr>
</thead>
<tbody>
<tr>
<td>2001</td>
<td>78</td>
<td>33</td>
<td>20</td>
<td>174</td>
</tr>
<tr>
<td>2002</td>
<td>73</td>
<td>19</td>
<td>23</td>
<td>198</td>
</tr>
<tr>
<td>2003</td>
<td>79</td>
<td>67</td>
<td>13</td>
<td>197</td>
</tr>
<tr>
<td>2004</td>
<td>97</td>
<td>57</td>
<td>21</td>
<td>181</td>
</tr>
<tr>
<td>2005</td>
<td>91</td>
<td>107</td>
<td>32</td>
<td>201</td>
</tr>
</tbody>
</table>
In the second model, we added hydrographical covariates to the geographical model, namely salinity, geostrophic velocity, and an index of water temperature. This model will be referred as the ‘hydrographical model.’ The selections of relevant hydrographical covariates were in part based on previous univariate analyses (Alemany et al. 2010) and in part based on minimizing cross-correlation among intervening variables. Salinity (Sal) was chosen because it is considered to be the best descriptor of the origin of water masses in the area (López-Jurado et al. 2008). Average salinities <37.5 describe inflows of fresh Atlantic waters from the southern areas, whereas typical resident Atlantic waters described by average salinities >37.5 are usually located northwards (Fig. 1). The geostrophic velocity (Gvel) was included in the model since it is positively correlated with the intensity of the fronts due to the confluence of fresher and older Atlantic water in the channels. Because the temperature in the mixed layer (Temp) was strongly correlated with the sampling date, we estimated the temperature residuals against day of the year and position (lat and lon) using a GAM. As such, positive (negative) temperature residuals indicate locations where water temperature is above (below) average, given the location and time of the year.

The hydrographical model was formulated as:

\[
\text{Stage1}_{\text{species}} = a_y + \text{offset} (\log (\text{vol})) + \text{factor(year)} + s_1 (\text{lat}, \text{long}) + s_2 (\text{cd}) + s_3 (\text{hour}) + s_4 (\text{Temp}) + s_5 (\text{Sal}) + s_6 (\text{Gvel})
\]

For both models, the variable selection criteria were based on the confidence region for the smooth effect, the R^2, and the generalized cross validation (GCV) score (Wood 2006). GAMs were fitted using the ‘mgcv’ library in R statistical software (www.r-project.org).

**RESULTS**

We determined 3 spatially and environmentally separated spawning areas. Bullet tuna mostly spawn inshore in coastal areas surrounding the island of Mallorca, whereas albacore and bluefin tuna spawn offshore in eastern and southern areas, respectively (Fig. 2). Bullet tuna was the species most consistently found in specific geographic locations, followed by albacore and bluefin tuna, the latter of which showed the most variable spatial distribution (Table 1, Figs. 3 to 5). The spatial patterns were well captured by our best models in the 3 species for every year (Figs. 3 to 5). The spawning activity (as detected by the stage-1 larval abundance) had shorter duration for bluefin tuna than albacore and bullet tuna, whose duration was more protracted (Fig. 6a−c). The time of the day effect showed that the catchability of all tuna species increased during the night, but bluefin tuna also showed a peak around midday (Table 2, Fig. 6d−f).

Oceanographic conditions strongly constrained spawning of bluefin tuna compared to albacore and bullet tuna, whose spawning habitat was mostly...
Reglero et al.: Spawning distributions of tuna in the Mediterranean

related to the spatial position (Table 2). This result is inferred from the changes in percentage of explained variance after inclusion of hydrographical variables in the geographical model. For bluefin tuna, the explained variance changed from 37.3% in the geographical model (of which <3% was explained by position alone, Table 2) to 75.3% in the total model. In contrast, the other 2 species had more contained changes of explained variance between geographic and total models (15.4 to 18.1% in albacore and 18.6 to 25.7% in bullet tuna). Bluefin tuna spawned in areas with intermediate salinities, where new and resident Atlantic waters converge, with a preference for areas with high frontal activity as shown from the positive relationship between bluefin tuna larval abundances and geostrophic velocity (Table 2, Fig. 7a−c). Temperature residuals had a significant effect only for bluefin tuna, with greater larval abundances at colder temperatures for a given location and time of the year. The best model fit for albacore was obtained for intermediate salinity and slow and intermediate geostrophic velocities, although the model improved little compared to the results obtained considering only geography (Table 2, Fig. 7d,e). For bullet tuna, there was a preference for low salinities and a positive linear relationship between the abundance of larvae and high geostrophic velocities, but still little improvement compared to the geographical model (Table 2, Fig. 7f,g). The distributions of the larval abundance for the 3 species in relation to background salinity and geostrophic velocities for 2001 to 2005 are shown in the supplement at www.int-res.com/articles/suppl/m463p273_supp.pdf.

DISCUSSION

Our results indicated that aggregations of tuna species spawn in the same large-scale area, but segregate their habitats at the regional scale based on the prevalence of either environmental or geographical preferences for their spawning habi-
Therefore, spawning adaptation to changing environments and the mechanisms to maximize fitness may differ among tuna species. Descriptions of spawning habitats for tuna have focused on single species, mostly bluefin tuna, e.g. western *Thunnus thynnus* in the Gulf of Mexico (Teo et al. 2007, Muhling et al. 2010) and *T. orientalis* in Japan (Kimura et al. 2010, Kitagawa et al. 2010). Only 1 study compared habitat characteristics among tuna species using univariate models (Alemany et al. 2010). Our study built upon these previous ones by producing a comprehensive understanding of tuna spawning phenology and geography across phylogenetically close species and over different environmental variables. The species' life history traits may explain the observed temporal and spatial portioning of the spawning habitat at the regional scale. A conceptual representation of how the 3 different species of tuna that we targeted in our analysis split their spawning habitats is shown in Fig. 8 and is further discussed below.

**Portioning of temporal spawning**

We observed that bluefin tuna had a confined temporal schedule for spawning compared to albacore and bullet tuna. The prevalence of the time of spawning of bluefin tuna from mid-June to early July in the Balearic Islands has been well documented (Medina et al. 2002, Corriero et al. 2003). Albacore and bullet tuna can still occur in high densities later in the season (e.g. August) when bluefin tuna larvae are scarce (Alemany et al. 2006, Torres et al. 2011). This reproductive trait probably relates to the species-specific thermal tolerances with regard to spawning and larval survival. In general, tuna larvae have a narrower and warmer range of temperature preferences than the adults (Boyce et al. 2008), in turn suggesting that spawning habitats are more restricted than adult foraging grounds. Bluefin tuna follow a similar reproductive strategy (i.e. confined seasonal spawning) in different spawning regions (Kimura et
In comparison, albacore and bullet tuna can spawn year-round in tropical waters (Klawe 1963, Ahlstrom 1971, Nishikawa et al. 1985, Ramon & Bailey 1996). No research has been conducted on the existing differences, if any, in the thermal tolerances of the larvae. Thus it is difficult to infer the degree to which physiological limitations determine the timing of spawning during the early life stages of the 3 species. Nevertheless, data from other temperate spawning areas support a wider temperature range for bullet tuna larvae than the other tuna species (Boehlert & Mundy 1994, Muhling et al. 2010). The tropical nature of bullet tuna and albacore is reflected in our analyses, since the effect of temperature residuals was only significant for bluefin tuna, with a preference to spawn in the colder than average water temperatures; this result fits well with bluefin tuna adults, which are less tolerant of warm waters (Blank et al. 2004).

Adaptive behaviors for spawning time may act as regulators of other processes such as larval trophic interactions. The displacement of the spawning window of bullet tuna and albacore could be an adaptation to reduce the larval trophic interactions, thus increasing offspring success. Especially in low-productivity environments, larval growth and survival depend on maintaining a pure zooplankton diet, later changing to a piscivorous and cannibalistic diet (Reglero et al. 2011). If inter- and intraspecific predation among larvae are important, then bullet tuna and albacore could have evolved a more protracted spawning timing to avoid larval predation and to increase offspring survival.

Lastly, the confined temporal window of bluefin tuna could be an adaptation for the adults that spend energy migrating large distances from optimal foraging grounds in the Atlantic Sea. Bluefin tuna most probably use stored energy to develop the gonads (Medina et al. 2002). Their spawning events are sporadic, and the adults stay in the spawning ground no more than 2 wk, leaving soon thereafter (Gunn & Block 2001, Medina et al. 2007).
Adult eastern bluefin tuna follow the Atlantic waters to spawn in the Mediterranean, which explains the prevalence of the recently hatched larvae in the southern area of the islands where the newest Atlantic water encounters the more saline resident surface water. The other 2 main spawning areas in the Western-Central Mediterranean are located around Malta and the South Tyrrenian Sea (Rooker et al. 2007), all receiving Atlantic water (Fig. 1). In the Balearic Islands, the position and activity of the fronts may act as a natural boundary that could be used by bluefin tuna adults as a signal to identify favorable spawning locations. Larger abundances of newly hatched albacore larvae in the northeast side of the Balearic Islands indicates spawning in surface waters with salinities above 37 and up to the maximum range, indicative of Atlantic water that has remained longer in the Mediterranean. For albacore, the dominant variable defining their spawning habitat was geographical position,
Fig. 7. _Thunnus thynnus_, _T. alalunga_, and _Auxis rochei_. Partial effects of significant environmental effects in the mixed layer (salinity, geostrophic velocity and temperature residual) on species abundance, as estimated from the hydrographical generalized additive model: (a–c) bluefin, (d,e) albacore and (f,g) bullet tuna. Other details as in Fig. 6.

Fig. 8. _Thunnus thynnus_, _T. alalunga_, and _Auxis rochei_. Conceptual figure summarizing our results, showing the spawning geography of the 3 species of tuna associated with their plausible migration pattern, for the large migratory bluefin tuna (black solid line), the Mediterranean albacore (grey solid line), and the coastal bullet tuna (black dotted line). Tuna pictures from S. P. Iglésias.
resulting in consistent hotspots of larvae in locations to the north-east of the Balearic Islands. This area is situated northwards of a consistent hydrographic feature, viz. a topographically induced eddy associated with the islands. The eddy occurs every summer (J. L. López-Jurado pers. obs.) and could be used as a predictable signal for spawning. The more coastal location of adult bullet tuna (Sabatés & Recasens 2001) fits well with the preference for coastal spawning areas. In contrast to albacore and bluefin tuna, large numbers of bullet tuna are also commonly found in coastal areas of the mainland, far from the islands (Morote et al. 2008), suggesting that spawning in this species is ubiquitously distributed in coastal areas.

The 3 species are iteroparous (spawning in several batches), asynchronous (having oocytes in all developmental stages), and indeterminate (new oocytes can develop in the female and be released during spawning) (Medina et al. 2002, Murua & Saborido-Rey 2003). The absolute fecundity is much lower in bullet tuna and albacore than in bluefin tuna due to their smaller size of the former, even though their relative batch fecundity is similar, around 59 eggs g\(^{-1}\) (Medina et al. 2007, Saber et al. 2011). Because of its large size, bluefin tuna can also store larger amounts of energy for reproduction and tolerate a broader range of environmental scenarios (Sharp 2001, Medina et al. 2002) than the other 2 species. This favors an opportunistic spawning behavior following specific environmental signals indicating that conditions are optimal.

Environmentally-driven spawning has been observed in bluefin tuna in other areas at a regional scale. Atlantic bluefin tuna belonging to the so-called Western stock migrate long distances to breed close to mesoscale structures in the Gulf of Mexico (Teo et al. 2007). On the other hand, short distance migrants or species with limited home ranges are limited in their opportunities to disperse during early life stages, since water parcels with retention features are typically located near bathymetric features (e.g. banks, capes, sills in fjords). This strategy will ensure self recruitment, probably at a more regional scale in albacore, spawning offshore in geographically fixed areas driven by predictable and consistent oceanographic features, and at a more local scale in bullet tuna, spawning adjacent to the coast. In Mallorca, the coastal morphology and wind forcing favors local recruitment of fish species inhabiting very coastal habitats (Basterretxea et al. 2012), a mechanism that may apply to bullet tuna as well at a larger spatial scale.

### Spawning strategies and capacities for adaptation to changing environments

Species responding to environmental cues shift their spawning distribution over contrasting environmental regimes, while species that respond to geographical cues will be less likely to change their spawning distribution in the face of underlying environmental changes. Bluefin tuna are opportunistic spawners that follow environmental signals and adapt to year-to-year or longer-time variability. A major physiological constraint for bluefin tuna during spawning is the lower tolerance for warm waters, mainly in larger fish (Blank et al. 2004). This may become an important limitation if the average surface temperatures increase in the Mediterranean Sea. If the reproductive success of tuna in the area is related to the coexistence of different developmental stages of tuna species, then different phenological responses among the 3 species to the environment influencing their timing and spatial spawning may influence their recruitment. On the other hand, migration constraints for the smaller-sized bullet tuna and albacore will limit expansion to new spawning habitats if these species are faced with environments less suitable for their larvae. Better knowledge of larval physiology and migration patterns are key to improving our understanding of the capacity for adaptation to changing environments in these species.

### Implications for management

Differences among the 3 species in their spawning strategies can play an important role for tuna conservation and management in the region. Currently, the major commercial fishery of tuna in the Mediterranean Sea consists of purse seiners explicitly targeting aggregations of reproductive eastern bluefin tuna adults (Fromentin & Powers 2005). Reproductive albacore and bullet tuna are also caught as by-catch or are targeted by traditional and recreational fisheries (ICCAT 2011). During the last decades, there has been a decreasing trend in eastern bluefin tuna spawning stock biomass (ICCAT 2008). Since spawning adults are the bulk of the fishery, temporal and/or spatial management measures such as closures to protect spawners, in addition to total allowable catch quotas already implemented, could be used as a strategy to manage and restore tuna populations. The information herein on the species’ reproductive traits is fundamental for spawning area closures to be
implemented successfully, a management option that has been described as biologically and economically efficient in other tuna stocks (Armsworth et al. 2010). Nowcasts and short-term forecast operational decisions including our results, that provide insights into the spawning habitat distributions of tuna species, could be a valuable tool for tuna conservation in the region (Hobday et al. 2011).

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