Trophic signatures of small-scale fishing gears: implications for conservation and management

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ABSTRACT: We quantified the ecosystem effects of small-scale gears operating in southern European waters (Portugal, Spain, Greece), based on a widely accepted ecosystem measure and indicator, the trophic level (TL). We used data from experimental fishing trials during 1997 to 2000. We studied a wide range of gear types and sizes: (1) gill nets of 8 mesh sizes, ranging from 44 to 80 mm; (2) trammel nets of 9 inner panel mesh sizes, ranging from 40 to 140 mm; and (3) longlines of 8 hook sizes, ranging from Nos. 15 (small) to 5 (large). We used the number of species caught per TL class for constructing trophic signatures (i.e. cumulative TL distributions), and estimated the TL at 25, 50 and 75% cumulative frequency (TL25, TL50, TL75) and the slopes using the logistic function. We also estimated the mean weighted TL of the catches (TL_W). Our analyses showed that the TL characteristics of longlines varied much more than those of gill and trammel nets. The longlines of large hooks (Nos. 10, 9, 7, 5) were very TL selective, and their trophic signatures had very steep slopes, the highest mean TL50 values, very narrow mean TL25 to TL75 ranges and mean TL_W > 4. In addition, the mean number of TL classes exploited was smaller and the mean TL50 and TLW were larger for the longlines of small hooks (Nos. 15, 13, 12, 11) in Greek than in Portuguese waters. Trammel and gill nets caught more TL classes, and the mean slopes of their trophic signatures were significantly smaller than those of longlines as a group. In addition, the mean number of TL classes exploited, the mean TL50 and the TLW of gill nets were significantly smaller than those of trammel nets. We attribute the differences between longlines of small hooks to bait type, and the differences between all gear types to their characteristic species and size-selectivity patterns. Finally, we showed how the slope and the TL50 of the trophic signatures can be used to characterise different gears along the ecologically ‘unsustainable–sustainable’ continuum.

KEY WORDS: Trophic level · Trophic signatures · Gill nets · Trammel nets · Longlines · Indicators · Northeast Atlantic Ocean · Mediterranean Sea

INTRODUCTION

Fishing has dramatic direct and indirect impacts on both coastal and open-sea marine ecosystems. Its imprint is identifiable at different time scales and levels of biological organisation (i.e. populations, communities and ecosystems) (Jackson et al. 2001, Pauly et al. 2002, Stergiou 2002, Christensen et al. 2003, Myers & Worm 2003, Lotze & Milewski 2004). Present-day marine ecosystems are impoverished versions of their pristine counterparts in terms of both species diversity and biomass (e.g. Jackson et al. 2001, Baum et al. 2003, Christensen et al. 2003, Myers & Worm 2003, Lotze & Milewski 2004), and are thus altered with respect to
between- and within-species interactions (see e.g. Pauly et al. 1998, Steele & Schumacher 2000, Pauly & MacLean 2003). Although the detrimental effects of fishing became apparent as early as the 1900s, global, multi-level effects have attracted strong attention only recently (e.g. Pauly & Christensen 1995, Pauly et al. 1998, Jackson et al. 2001, Baum et al. 2003, Christensen et al. 2003, Myers & Worm 2003, Pauly & MacLean 2003; reviews by Botsford et al. 1997, Jennings & Kaiser 1998, Hall 1999, Pauly et al. 2002, 2003). The results of these studies alarmed scientists, international organisations, non-governmental organisations and the public at large, and raised the issue of placing fisheries resource management within an ‘ecosystem’ context (e.g. Brownman & Stergiou 2004). This led researchers to start exploring the use of a plethora of indicators in capturing/expressing fishing-induced effects at the community or ecosystem level. Among the various measures (e.g. Caddy & Garibaldi 2000, Pinnegar et al. 2002, Nicholson & Jennings 2004), those based on the fractional trophic level (TL), which expresses the position of an organism in the food web (Froese & Pauly 2000), enjoy wide acceptance (i.e. one measure, the mean trophic index, has been selected by the Conference of the Parties to the Convention of Biological Diversity as 1 of 8 biodiversity indicators: see Pauly & Watson 2005).

The strength of TL as an ecological indicator lies in its efficiency in capturing and revealing ecosystem effects of fishing, either: (1) directly, in the form of the mean weighted TL of the catch for a series of years, useful for identifying the ‘fishing down the marine food webs’ process, which was described for the first time by Pauly et al. (1998, 2001) and later applied by others (e.g. Vasconcelos & Gasalla 2001, Pinnegar et al. 2002, 2003, Nicholson & Jennings 2004), or (2) indirectly, for estimating other indicators such as the primary production required to support fisheries (Pauly & Christensen 1995; and applications in CIEM 2000, Vasconcelos & Gasalla 2001) and the ‘Fishery in Balance’ index (Pauly et al. 2000a). In addition, recently, McClanahan & Mangi (2004) have also used the mean weighted TL of the catches of different small-scale gears as a simple index for the management of Kenyan small-scale fisheries. The TL concept also opened up new topics for marine ecological research, such as: (1) the comparison of different ecosystems based on the TL frequency distribution of the relevant species (known as trophic signatures: Froese et al. 2005) and (2) the relationship between TL and other biological variables, notably size (between species: Froese & Pauly 2000; ontogenetic changes: Pauly et al. 2001, Stergiou & Karpouzi 2002) and mouth area (Karpouzi & Stergiou 2003).

Fishing is practised with a variety of gears, many of which exploit the same stocks or different parts of the same stock (e.g. Stergiou et al. 2004). Gear selectivity, both in terms of body sizes and species, differs with gear type, with gears typically ranging from unselective to highly selective. For example, trawls, purse seines and beach seines are relatively unselective in terms of species and sizes, characterised by logistic type size selectivity (Millar & Fryer 1999). On the other hand, gill nets are more size selective, largely showing unimodal size selectivity (Millar & Fryer 1999, Erzini et al. 2003). Compared to gill nets, trammel nets and longlines are generally less size selective, having multiple size-selectivity types (Erzini et al. 1996, 2003, Anonymous 2001, Stergiou & Erzini 2002). In addition, modern fishing vessels and technology nowadays allow fishers access to practically all fish habitats, which often differ in terms of the TL ranges of the inhabiting fishes (Stergiou & Karpouzi 2002). Thus, an important question is whether the number of species caught per TL class as well as the mean weighted TL of the catches differ with gear type, which would imply that the gears’ effects on the ecosystem differ accordingly.

In this report, we quantified the trophic impacts of small-scale fishing gears in southern European waters, which historically support very important small-scale fisheries (OCEANA 2004), using data from experimental fishing trials conducted during 1997 to 2000. To that end, we compared the cumulative TL frequencies (henceforth called trophic signature, based on Froese et al. 2005) as well as the mean weighted TL of the catches of a variety of small-scale fishing gears (i.e. trammel nets of 9 inner mesh sizes, gill nets of 8 mesh sizes and longlines of 8 hook sizes) in the Northeast Atlantic Ocean (Portugal: Algarve waters; Spain: Gulf of Cádiz) and the eastern Mediterranean Sea (Greece, Cyclades) (Fig. 1). In particular, we tested whether (1) the trophic signatures of small-scale gears differ with gear type, gear size and area, and (2) the mean weighted TL of the catch differs with gear type, gear size and area. Consequently, we propose how the statistical properties of the trophic signatures, notably the slope and the TL at 50% cumulative frequency, can be used as an indicator of the gear-specific effects at the ecosystem level and, thus, as a tool for the conservation and management of fisheries resources.

**MATERIALS AND METHODS**

**Sampling.** Experimental fishing trials with gill and trammel nets of different mesh sizes and with longlines of different hook sizes took place in the Algarve, the Gulf of Cádiz and the Cyclades (Fig. 1), at different depth ranges during 1997 to 2000 (Table 1). All gears were set on the bottom, targeting demersal species. In
all areas, sampling was conducted using commercial boats, and fishing operations were carried out by professional fishers who fished on traditional fishing grounds following their normal operating practices. Erzini et al. (1999, 2001, 2003), Anonymous (2001), Stergiou et al. (2002) and Stergiou & Erzini (2002) provide a thorough description of the fishing gears used and of the sampling schemes adopted in each region. To separate the catches and record the necessary data, 1 to 3 researchers were onboard. Weight per species was not recorded in the case of longlines of large hooks (Nos. 5 to 10, where hook No. 1 is the largest possible and hook No. 15 is among the smallest ones in commercial fisheries).

**Classification by trophic level.** Overall, 178 taxa were caught, of which 175 were identified to the species level (1 echinoderm, 7 crustacean, 6 cephalopod and 161 fish species) and 3 to genus, or higher level (fishes: Gobiidae, Labrus and Raja). For 65 of the 161 fish species we extracted TL values from Stergiou & Karpouzi (2002), who collected all published information on the feeding habits and estimated TL for 332 Mediterranean fish stocks, belonging to 146 species. For 84 fish species, we extracted TL values from FishBase online (www.fishbase.org), which provides TL estimates from food items and diet compositions (together with full documentation on TL estimation) for a large number of fish species. For the remaining 12 fish species, no TL estimate was available in FishBase. However, TL is rather phylogenetically constrained, being a function of species’ body length (Froese & Pauly 2000) and mouth area (Karpouzi & Stergiou 2003), with families often having distinct TL ranges (Stergiou & Karpouzi 2002). Thus, for these 12 species we searched FishBase for TL values of other species belonging to the same genus (using the FishBase valid genus of the species). We found TL for 8 species (i.e. Ariosoma balearicum, Microchirus azevia, Microchirus ocellatus, Peristedion cataphraetum, Raya brachyurops, Stephanocephalus diaspros, Synaptura lusitanica, Trachinus radiatus). For these 8 species, we used the mean TL value of all species per genus (with the number of species/genus ranging from 1 to 18 species). For the 4 fish species (i.e. Symphodus bailloni, Monochirus hispidus, Ophisurus serpens and Synapturichtys kleinii) for which no TL estimate was available for any other species of the genus, we used TL estimates from the ‘Tools/Ecopath data’ option of FishBase. Finally, for the 3 fish taxa identified to genus or higher level, we used the mean TL of all species of the genus in FishBase for Raja and Labrus and TL = 3.5 for Gobiidae (the default value used in TrophLab, a software used for the estimation of TL from diet data, downloadable from www.fishbase.org; Pauly et al. 2000b). The TL values of cephalopods and crustaceans were extracted from the Sea Around Us database (http://seaaroundus.org/catchrate/map/trophic_levels.htm), whereas for the 1 echinoderm taxon we used TL = 2.4 (the default TrophLab value).

**Trophic signatures.** Subsequently, we estimated the number of species per TL class (30 classes, TL range from 2 to 4.9, Step 0.1) for the 102 gear/size/season/area combinations and for the 29 gear/size/area combinations (i.e. all seasons combined per gear/size/area). We note that in the case of the large hooks in the Algarve, data refer to years (1997, 1998), not seasons. From the number of species per TL class we constructed cumulative TL frequency distributions (i.e. trophic signatures) for each case. Based on the characteristics identified from these 2 sets of trophic signatures, we also constructed ‘general’ trophic signatures for each gear (gill and trammel nets: all sizes/seasons/areas combined; longlines: all seasons combined by large and small hook sizes and area).
For comparative purposes we also extracted from FishBase (www.fishbase.org.ph; using the option: ‘Information by Country/Ecosystem/Island/Ecopath models’) all fish species for which TL is available in Portuguese waters (used as representative of the Algarve and Cádiz areas) and in Greek waters, henceforth called ‘ecosystems’. Overall, TL was available for 482 and 337 species, respectively, out of a total of 484 and 339 marine species reported by FishBase for these 2 countries.

We compared the trophic signatures of the gears with each other and with those of the ecosystems. We constructed trophic signatures first, including all taxa caught and then including only fishes. We tested for differences in the TL frequency distributions only for the general trophic signatures, using the Kolmogorov-Smirnov test (KS).

Because cumulative TL frequency curves are of a logistic type, we described the relationship between the cumulative frequency (CF) of a TL class and TL using the logistic function:

\[
CF = \frac{1}{1 + e^{-b(TL - TL_{50})}}
\]

where \(b\) is the slope of the curve and \(TL_{50}\) is the TL corresponding to the 50\% cumulative frequency. Fits were obtained using non-linear, least-squares regression (SAS 1988). We also estimated the cumulative TL range corresponding to 25 and 75\%, \(TL_{25}\) and \(TL_{75}\), of the cumulative frequencies:

### Table 1. Study gears and sampling details for the experimental fishing conducted in the 3 geographic areas

<table>
<thead>
<tr>
<th>Gear</th>
<th>Algarve</th>
<th>Gulf of Cádiz</th>
<th>Cyclades</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Trammel nets</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inner mesh size (stretched mm)</td>
<td>100, 120, 140</td>
<td>80, 90, 100</td>
<td>40, 48, 56</td>
</tr>
<tr>
<td>Outer mesh size (stretched, mm)</td>
<td>600, 800</td>
<td>300, 400</td>
<td>220/240, 240/260, 280/300</td>
</tr>
<tr>
<td>Number of gears used(a)</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Sampling frequency</td>
<td>Seasonal</td>
<td>Autumn, spring</td>
<td>Seasonal</td>
</tr>
<tr>
<td>Total number of hauls (for each gear per season)</td>
<td>40</td>
<td>60</td>
<td>41</td>
</tr>
<tr>
<td>Combinations used in this study(b)</td>
<td>12</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>Sampling depths (m)</td>
<td>15–100</td>
<td>10–30</td>
<td>10–80</td>
</tr>
<tr>
<td><strong>Gill nets</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mesh sizes (stretched, mm)</td>
<td>50, 60, 70, 80</td>
<td></td>
<td>44, 48, 52, 56</td>
</tr>
<tr>
<td>Number of gears used</td>
<td>4</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Sampling frequency</td>
<td>Seasonal</td>
<td></td>
<td>Seasonal</td>
</tr>
<tr>
<td>Total number of hauls (for each gear per season)</td>
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<td>42</td>
<td></td>
</tr>
<tr>
<td>Combinations used in this study(b)</td>
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<td>16</td>
<td></td>
</tr>
<tr>
<td>Sampling depths (m)</td>
<td>15–60</td>
<td></td>
<td>4–90</td>
</tr>
<tr>
<td><strong>Longlines of small hooks</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hook size</td>
<td>15, 13, 12, 11</td>
<td>15, 13, 12, 11</td>
<td></td>
</tr>
<tr>
<td>Hook type</td>
<td>Mustad round bend</td>
<td>Mustad round bend</td>
<td></td>
</tr>
<tr>
<td>Number of gears used</td>
<td>4</td>
<td>4</td>
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<tr>
<td>Sampling frequency</td>
<td>Seasonal</td>
<td></td>
<td>Seasonal</td>
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<tr>
<td>Total number of hauls (for each gear per season)</td>
<td>40</td>
<td>43</td>
<td></td>
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<tr>
<td>Combinations used in this study(b)</td>
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<td></td>
</tr>
<tr>
<td>Sampling depths (m)</td>
<td>15–60</td>
<td></td>
<td>4–90</td>
</tr>
<tr>
<td><strong>Longlines of large hooks</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Hook size</td>
<td>10, 9, 7, 5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hook type</td>
<td>SIAPAL brand</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of gears used</td>
<td>4</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Sampling frequency</td>
<td>Annual</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total number of hauls (for each gear per year)</td>
<td>20</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Combinations used in this study(c)</td>
<td>8</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Sampling depths (m)</td>
<td>200–700</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Total combinations</strong></td>
<td>52</td>
<td>6</td>
<td>44</td>
</tr>
</tbody>
</table>

\(a\)The different (inner \(\times\) outer) mesh size combinations were combined based on inner mesh size because outer mesh size did not affect significantly species catch rates and size selectivity (Anonymous 2001)

\(b\)(sampling frequency) \(\times\) (number of gears used); all trials for each gear per season were combined

\(c\)(sampling frequency) \(\times\) (number of gears used) \(\times\) (2 yr); all trials for each gear per year were combined
\[ TL_{25} = \left( S_1 - \ln 3 \right) / S_2 \]

and

\[ TL_{75} = \frac{\ln 3 + S_1}{S_2} \]

where \( S_2 \) equals \( b \) and \( S_1 \) equals \( b(TL_{50}) \).

**Mean weighted trophic level.** We also estimated the mean TL of the catches by gear size, season (year in the case of the longlines of large hooks) and area weighted (\( TL_w \)) by the weight of each species participating in the catches. In the absence of weight measurements for longlines with large hooks, weighting was based on the numbers of each species caught, a fact that does not greatly affect the results given that the vast majority of specimens caught were of relatively large size.

We compared the mean \( TL_w \) of the catches across gears and areas using a \( t \)-test or 1-way ANOVA and Fisher’s least significance difference test. We explored the relationships between gear size and \( TL_w \) using regression analysis.

**RESULTS**

**Trophic signatures**

The seasonal trophic signatures of all trammel and gill net combinations were bounded in a narrower band when compared to those of longlines (Fig. 2a). They were evenly dispersed around those signatures representing the ecosystems for cumulative frequencies <50 to 60%, and to the left of those of the ecosystems at frequencies >50 to 60% (Fig. 2a). The trophic signatures of gill nets were shifted more to the left than those of the trammel nets (Fig. 2a). In contrast, the trophic signatures of the longlines were distributed in a very wide band around the ecosystem signatures (Fig. 2a). This was because of the different positions of the Algarve and Cyclades longlines of small hooks (Nos. 15 to 11; to the left and to the right of the ecosystem signatures, respectively), and of the Algarve longlines of large hooks (Nos. 10 to 5; further to the right of those from the Cyclades).

The same pattern emerged when seasonal (Fig. 2b) and size/area (Fig. 2c) effects were removed. In the last case (Fig. 2c), however, the difference between gill and trammel nets was largely smoothed out. Indeed, the trophic signature of all trammel nets combined (Fig. 2c) did not differ significantly (KS, \( p > 0.05 \)) from that of gill nets. In contrast, the trophic signatures of the longlines were again distributed in a wide band, with those of small and large hooks in Algarve waters...
were shifted to the far left and to the far right of the ecosystem signatures; those of small hooks in the Cyclades were between those of the Algarve (Fig. 2b, c). The TL frequency distributions of the longlines of small and large hooks in the Algarve and Cyclades (Fig. 2c) differed significantly (KS, p < 0.05) from each other, as well as from those of the trammel and gill nets. Finally, the TL frequency distributions of all gears (Fig. 2c) differed significantly (KS, p < 0.05) from those of the 2 ecosystems, whereas the latter did not differ from each other (KS, p > 0.05). The same pattern emerged when we considered only fish species in the analysis (graphs not shown).

The mean number of TL classes exploited per gear differed significantly (ANOVA, p < 0.05) between gears, being highest for trammel nets and lowest for longlines of large hooks in the Algarve and of small hooks in the Cyclades (Fig. 3). All gears exploited fewer TL classes than exist in the 2 ecosystems (n = 26).

The results of the application of the logistic curve to the trophic signatures of the 102 gear/size/area combinations showed that longlines as a group were characterised by the highest variation in slope, TL50 and TL25 to TL75 range (Fig. 4). The mean slope, TL50 and TL25 to TL75 range all differed significantly (ANOVA, p < 0.05) with gear (Fig. 4). The mean slope was highest for all longlines and smallest for gill and trammel nets (Fig. 4a). The mean TL50 was smallest for the longlines of small hooks in the Algarve and highest for the longlines of large hooks (Fig. 4b). Finally, the mean TL25 to TL75 range was narrower for all longlines than for gill and trammel nets (Fig. 4c).

Mean weighted trophic level

The TLW per gear size and season ranged between 3.09 and 3.92 for gill nets (Fig. 5a), 3.47 and 3.94 for trammel nets (Fig. 5b) and 3.18 and 4.12 for longlines (small hooks, Algarve: 3.33 to 3.45; small hooks, Cyclades: 3.18 to 3.72; large hooks: 4.07 to 4.12) (Fig. 5c). The mean TLW of gill nets (Fig. 5a) and longlines of small hooks (Fig. 5c) both differed significantly (t-test, p < 0.05) between Algarve and Cyclades. The mean TLW of trammel nets also differed significantly (ANOVA, p < 0.05) with area (Fig. 5b). With the exception of the gill nets in Algarve waters (Fig. 5a), where TLW was a square function of mesh size, no significant area-specific relationship was found between TLW and gear size.
for all gears (Fig. 5). However, for gill nets in the Cyclades (Fig. 5a) and trammel nets in the Algarve and Cyclades (Fig. 5b), the pattern indicated an increase in $TL_W$ with gear size. In contrast, for longlines there was clearly no relationship between $TL_W$ and hook size (Fig. 5c).

Finally, the mean $TL_W$ also differed significantly between gears (ANOVA, $p < 0.05$), being highest for longlines of large hooks and lowest for gill nets and longlines of small hooks in the Algarve (Fig. 6).

**DISCUSSION**

Small-scale fisheries, which account for about half of the fisheries biomass directly consumed by humans (FAO 2004), are more important, both economically and socially, than industrial and large-scale fisheries (Pauly & MacLean 2003). Thus, for instance, small-scale fisheries in Greek waters account for >90% of the total number of fishing boats, >65% of the total number of fishers and >50% of the total wholesale value of the Greek catch (Stergiou et al. 1997). Although industrial and large-scale fisheries are generally favoured by our political systems, the redistribution of effort from industrial and large-scale to small-scale fisheries would have important benefits for the socio-economy of the fisheries sector and the ecosystems themselves (Pauly & MacLean 2003). In this light, the assessment of the impact of the 2 types of fisheries at the community and ecosystem levels is crucial.

In this study we quantified the ecosystem effects of a large number of the most important small-scale gear sizes and types operating in southern European waters. Our quantification was based on a widely accepted ecosystem measure and indicator, notably TL (see 'Introduction'). The gears studied support very important fisheries in the 3 study areas (Erzini et al. 1996, 1999, 2003, Stergiou et al. 1997, 2002, Anonymous 2001, Stergiou & Erzini 2002) as well as in other southern European and Mediterranean countries (e.g. OCEANA 2004).

**Differences among gears**

Our analyses indicate that the different small-scale gears generally exploit fewer TL classes and in different proportions than those occurring in the ecosystem, and that the TL characteristics of longlines varied much more than those of gill and trammel nets. Our analysis also demonstrated that the trophic signatures of longlines of large hooks (baited with sardine *Sardina pilchardus*) differ from those of the remaining gears. The longlines of large hooks exploited the fewest TL classes, at high TL, and at proportions much higher than those occurring in the ecosystems. Their catch was dominated...
by carnivorous species, especially hake *Merluccius merluccius* and elasmobranchs (Erzini et al. 2001), with those exhibiting preference for fish and cephalopod prey making up about 50% of the catch (Fig. 7). As a result, their TLW was >4. These features were efficiently captured by the statistical properties of the trophic signatures. Thus, longlines of large hooks had very steep slopes, the highest mean TL50 values and a very narrow TL25 to TL75 range, being positioned to the right of the trophic signatures of the remaining gears and of the ecosystems.

Trophic signatures of longlines of small hooks differed in the Algarve and Cyclades. Thus, although the mean slope and TL25 to TL75 range of the longlines of small hooks in the Cyclades did not differ significantly from those in Algarve waters, the mean number of trophic level classes exploited was smaller and their mean TL50 and TLW values were larger than those in the Algarve. We attribute this difference to the different types of bait used in the 2 areas. In Algarve waters, the small hooks were baited with the worm *Sipuncula nudens* and thus caught more omnivores with preference for animal prey than in the Cyclades (Fig. 7). In the latter area, small hooks were baited with squid, and thus their catch included carnivores with a preference for cephalopods and fishes in higher percentages than in the Algarve (Fig. 7). Indeed, species selectivity regarding hooks depends largely on bait type (Bjordal & Lokkeborg 1996).

Trammel nets differed from gill nets with respect to all measures used in this study. Gill nets caught a somewhat greater number of omnivores with preference for animal prey, and fewer carnivores than did trammel nets (Fig. 7).

Accordingly, the gill net TLW was smaller than that of trammel nets and was comparable to that of the longlines of small hooks in the Algarve. In contrast, the mean TLW of trammel nets was the highest after that of the longlines of large hooks (Fig. 7). We attribute this difference to the fact that trammel nets are in general much less selective in terms of both sizes and species than gill nets, exhibiting a wide range of selectivity types, from unimodal to multimodal and logistic (Anonymous 2001, K. Erzini unpubl. data). As a result, trammel nets catch a wider size range, both within- and between-species, and, thus, because TL generally increases with size (Froese & Pauly 2000), a wider range of TL classes than gill nets.

Given that the biomass of large-sized species, generally positioned high in the food web (Froese & Pauly 2000), declines more rapidly than that of their small-sized counterparts (see discussion in Pauly et al. 2001), longlines of large hooks and small hooks baited with squid strongly contribute to ‘fishing down the marine food web’ by selectively removing high TL species. Catch and effort data from pelagic longlines of very large hooks from various areas of the world ocean show that biomass declines of large predatory fishes can be very rapid (i.e. as high as 80% in the first 15 yr: Baam et al. 2003, Myers & Worm 2003).

McClanahan & Mangi (2004) also compared the TLW of different small-scale gears in Kenyan waters. They found that the hand-lines of small hooks, usually baited with worms, pieces of crab, squid, or octopus, have sig-

![Fig. 6. Box-whisker plots of the mean TLW (mean weighted trophic level) of the catches. See Fig. 3 for definition of boxes and abbreviations.](image1)

![Fig. 7. Percentages of the different trophic functional groups (based on the species’ trophic levels) participating in the catches of gill nets, trammel nets, longlines of small hooks in the Algarve (A), longlines of small hooks in Cyclades (G) and longlines of large hooks in the Algarve. The number ranges correspond to the TLW ranges shown in Fig. 5. The numbers in parentheses correspond to the mean TLW shown in Fig. 6. Trophic functional groups and their trophic level ranges, in parentheses, are from Stergiou & Karpouzi (2002), who estimated trophic levels from diet compositions for 332 Mediterranean fish stocks belonging to 146 fish species.](image2)
nificantly higher TL$_{W}$: 3.6, than gill nets of 60 mm, beach seines, spears and big and small traps, all of which had TL$_{W}$ values of between 2.6 and 2.9. These authors also claimed that hand-lines, by targeting the large-sized high-trophic-level species such as snappers, emperors and wrasses, contribute to 'fishing down the marine food web' in Kenyan waters. It is interesting to note that the TL$_{W}$ of the Kenyan hand-lines is comparable to that of the longlines of small hooks in the Cyclades.

With the exception of the Algarve gill nets, no significant relationship was found between gear size and TL$_{W}$, although there was some evidence of an increase in TL$_{W}$ with mesh size for gill nets and, to a lesser extent, for trammel nets. This might be the result of the fact that in the present study we did not account for ontogenetic changes in TL. Indeed many studies show that the TL of many species, especially those of omnivores with preference for animal prey and carnivores, increases asymptotically with size during their life (e.g. stomach contents: Stergiou & Karpouzi 2002; isotope analysis: Badalamenti et al. 2002). Thus, one would expect that TL will also increase with gear size. This is especially true for gill nets, for which larger mesh sizes generally catch larger specimens of the same species, and less so for trammel nets and longlines, where catch distributions for different mesh and hook sizes tend to overlap (Anonymous 2001, Stergiou & Erzini 2002, Erzini et al. 2003).

The unexpected square relationship in the case of the Algarve gill nets might be the result of the exceptionally high catches of Scomber japonicus in the 80 mm gill nets (35% of the total catch by weight; Erzini et al. 1999). Its TL of 3.1 is much lower than the TLs of the species that dominated the catches of the 50 and 60 mm gill nets (i.e. Scorpaena notata, Mullus surmuletus and Diplodus bellottii: TLs between 3.4 and 3.6).

**Ecosystem effects**

Overall, the statistical properties of the trophic signatures, i.e. slope and TL$_{50}$, captured the characteristics of the different gears in terms of exploited TL classes and, thus, of their impact on the ecosystem. Therefore, the gear-specific TL$_{50}$–slope combination of a trophic signature can be used as an indicator for managing fisheries at the ecosystem level. In general, an indicator is useful when it is easily understood and estimated, and can be attached to both a reference point and an objective (Gislason et al. 2000). Here, the objective is the trophic level balance of the ecosystem, and the reference point is a specific set of TL$_{50}$–slope values that can be related to ‘ecologically sustainable fishing’. When complete knowledge exists of all metiers (i.e. combinations of specific gear, gear size, season and area target-

ing a species or group of species: sensu Ulrich et al. 2001) operating in an ecosystem, and of their catch species compositions, then categorising/selecting metiers in terms of ‘sustainable’ trophic signatures is an attainable goal. We illustrate this in Fig. 8, in which TL$_{50}$ is plotted against the slope for 3 hypothetical gears (shown as an inset) as well as for the 102 cases used in the present study. The 3 hypothetical gears include: (1) 1 gear exploiting only the very low TL classes (e.g. a bivalve dredge); (2) 1 gear exploiting only the highest TL classes, such as for instance the longlines of the largest hook sizes that catch large pelagics and sharks; and (3) 1 gear exploiting low and high TL classes at higher and lower percentages, respectively, than those in the ecosystem, having a trophic signature of moderate, almost linear slope (e.g. a rather unrealistic gear, roughly approximated by a purse seine catching mainly small- and medium-sized pelagic organisms, but also other demersal and bentho-pelagic species of high TL).

In such a plot, the vertical and horizontal lines intersecting at the points defined by the TL$_{50}$–slope combination of the corresponding ecosystem define 4 areas of TL$_{50}$–slope combinations (A to D: Fig. 8). No TL$_{50}$–slope combination is possible for Area D, whereas gears with combinations falling in Area A are also very rare. Thus, the vast majority of gears will have trophic signatures with TL$_{50}$–slope combinations falling either in Area B or C, and their trophic characteristics (summarised in Fig. 8) will increase along the diagonals. We propose that fisheries based on gears with TL$_{50}$–slope combinations positioned within the shaded area in Fig. 8, i.e. around those of the ecosystem, lie towards the ecologically ‘sustainable’ end of the ‘unsustainable–sustainable’ continuum. This is because these gears are less TL selective and generally remove low and high TL classes in percentages higher and lower, respectively, than those found in the ecosystem. Examples of such ‘sustainable’ gears are all trammel net combinations, as well as the hypothetical Gear 3 (Fig. 8), provided that they are not excessively used. In contrast, fisheries based on gears with TL$_{50}$–slope combinations falling outside this shaded area are towards the ecologically ‘unsustainable’ end, especially those based on gears positioned further away along the diagonals (Fig. 8). Examples of such gears are: (a) all longlines of large hooks and the hypothetical Gears 1 and 2 and (b) the Cyclades longlines of Hook 11 in winter, which largely caught species of TL values >3.7, and of Hook 13 in spring, which largely caught species of 1 TL class, 3.7 to 3.8, corresponding to the uppermost and rightmost filled circles, respectively (Fig. 8). Gears falling outside the shaded area in B remove high TL classes at proportions much greater than those existing in the ecosystem, thus contributing to ‘fishing down the marine food web’. Fisheries based on these gears can
drive top-down cascade effects, which are common in marine ecosystems (e.g. coastal areas: Jackson et al. 2001; open oceans: Myers & Worm 2003; enclosed seas: Daskalov 2002), with important consequences for their structure and dynamics. In contrast, gears falling outside the shaded area in C remove low TL classes at proportions much higher than those that exist in the ecosystem, a fact that might drive bottom-up cascading effects (see e.g. Cury et al. 2000). Thus, the use of both types of gears should be reduced.

Limitations

Our study, which is the first one to propose the use of trophic signatures in capturing the ecosystem effects of small-scale gears, suffers from limitations, which refer to both the data and the approach. With respect to the data used, our study was based on TLs estimated from diet data. This method, as all methods used for estimating TL, suffers from certain disadvantages (for a discussion see CIESM 2000, Polunin & Pinnegar 2002). In addition, the TL of some species, such as those of the genera Liza and Mullus, are difficult to determine (see e.g. Pinnegar et al. 2003). Thus, TL estimates are uncertain. Ideally, even in the case of accurately estimated TLs, one should use area–year–length-specific TL per species rather than ‘global’ or ‘rounded’ ones in order to minimise uncertainty. At present, however, high-resolution TL data are not available for most marine animal species. We minimised the effect of data ‘accuracy’ by using either TL estimates from the Mediterranean (for 65 species: from Stergiou & Karpouzi 2002) or ‘average’ species-specific TLs (i.e. referring to the species in general: for 97 species) whenever possible. Thus, ‘rounded’ or ‘global’ TL values were used in only 16 of 178 taxa. In our approach, ecosystem trophic signatures have a central role: they are used to define the 4 rectangles in Area D, as well as for those analysed in the present study (102 cases) (filled rhombus: the 2 ecosystems [where the 2 lines intersect]; filled circles: longlines of small hooks; open circles: longlines of large hooks in Algarve; open triangles: trammel nets; open rhombus: gill nets). The main characteristics of gears with TL_{50–slope} combinations positioned in different areas (A to C) are also shown. For explanation of the shaded area and arrows see ‘Discussion’.

Fig. 8. The relationship between trophic level at 50% cumulative frequency (TL_{50}) and slope of the trophic signatures of 3 hypothetical gears (indicated by the 3 large filled squares, 1 to 3), shown as an inset in Area D, as well as for those analysed in the present study (102 cases) (filled rhombus: the 2 ecosystems [where the 2 lines intersect]; filled circles: longlines of small hooks; open circles: longlines of large hooks in Algarve; open triangles: trammel nets; open rhombus: gill nets). The main characteristics of gears with TL_{50–slope} combinations positioned in different areas (A to C) are also shown. For explanation of the shaded area and arrows see ‘Discussion’.
clearly predominate at TL >3 (Froese et al. 2005). Indeed, trophic signatures were exactly the same after the exclusion of the non-fish taxa, one of which dominated catch species composition of the trammel nets (i.e. Sepia officinalis) in the Algarve and the Gulf of Cádiz: Anonymous 2001). Yet we preferred to include rather than exclude non-fish taxa, in order to capture the actual gear-specific trophic signatures.

**Outlook**

The above-mentioned limitations affect the practical applicability rather than the performance of the approach itself. The TL concept has attained a recognised importance (i.e. see ‘Introduction’). Thus, we believe that in the near future increased research effort will be devoted to obtaining reliable region- and length-specific TL estimates for most marine animals, either from diet or stable isotope data.

The trophic signature approach gives the same weight to both locally abundant species, for which good TL estimates are usually available, and to locally rare species, for which less accurate TL estimates are usually available. Many studies describing long-term temporal trends in the mean TL of landings (e.g. Pauly et al. 1998, 2001, Vasconcelos & Gasalla 2001, McClanahan & Mangi 2004) are also based on diet-estimated TLs. However, in such cases this is less of a problem, because the mean TL is biomass weighted. In the current study, the fact that the results of the TLW analysis agreed with those of the non-weighted trophic signature approach (Fig. 7) is an indication that the lack of weighting might not be a real impediment.

Finally, the proposed shading in Fig. 8, which indicates the area of ‘sustainable’ TL50–slope combinations, is subjective. Yet a more objective delineation of the shape and dimension of this area is critical for the definition of reference points. This, however, can be realised only after various issues or factors have been accounted for, such as: (1) full knowledge of all metiers and of the ecosystem in which these metiers operate; (2) application of this approach to fisheries at different maturity stages, since the latter affect the gear-specific, catch-species composition (McClanahan & Mangi 2004); and (3) application of this approach in other areas, covering a variety of depth ranges, habitats, fishing gears and gear sizes. Accounting for such issues will eventually allow the sustainable harvest from a particular ecosystem using several fishing gears, each targeting different TL classes (i.e. having different trophic signature). Naturally, this implies that TL harvest rates must match the ecosystem’s capacity to replenish the biomass removed from the impacted TL classes. Thus, further empirical and theoretical studies are needed to define the extent of this reference area and the practical applicability of our approach to conservation and management of fisheries resources.

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