



Spillover from six western Mediterranean marine protected areas: evidence from artisanal fisheries

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ABSTRACT: This study investigated spillover (biomass export) around 6 marine protected areas (MPAs) in the western Mediterranean based on catch and effort data from artisanal fisheries. The selected MPAs were Cerbère-Banyuls and Carry-le-Rouet in France, and Medes, Cabrera, Tabarca, and Cabo de Palos in Spain. These MPAs had been functional for more than 8 yr and incorporate areas of fisheries closure and restricted use where fishing is limited. We based our study on the hypotheses that, in the presence of biomass export, (1) fishing effort would concentrate close to MPA boundaries, and (2) fishery production, expressed as catch per unit area (CPUA), would be highest near MPA boundaries and decrease with distance. We selected data from 14 'fishing tactics' using gill nets, trammel nets and bottom long-lines targeting sparids, mullids, serranids, scorpaenids and palinurids. We analyzed the spatial distribution of effort, fishery production and revenues per unit area, using generalized additive models (GAMs), and we tested regression slopes of effort density and CPUA with distance to closure boundaries, using generalized linear models (GLMs). GAMs allowed us to recognize habitat discontinuities or 'hot spots' of high production in the vicinity of the MPAs, and to identify the extent of potential spillover effects in order to implement GLMs. We found evidence of effort concentration and high fishery production near fisheries closures for all fishing tactics analyzed and significant negative slopes for most. Revenues generally followed trends similar to CPUA. Significant negative slopes from GLM of effort density and CPUA with distance from fisheries closures were indicative of biomass export where habitats across closure boundaries had some degree of continuity. The spatial extent of spillover was consistent with species mobility and fisheries efficiency and extended 700 to 2500 m from fishery closure boundaries. Our results suggest that coastal MPAs can be an effective management tool for artisanal fisheries in the region and can be extended to the rest of the western Mediterranean, as the fishing tactics studied are typical of the region.

KEY WORDS: Marine protected areas · Fisheries closures · Artisanal fisheries · Spillover · Fishing the line · Catch gradients · Mediterranean Sea

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INTRODUCTION

Fishing restrictions in marine protected areas (MPAs) can promote increases in biomass of exploited species, which under particular conditions will lead to spillover

(i.e. biomass export) to surrounding non-protected areas, maintaining or enhancing local fisheries (e.g. reviews in Roberts & Polunin 1991, Sánchez-Lizaso et al. 2000, Russ 2002). Species that have shown strong responses to protection are those that have moderate vagility in relation

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to the size of the protected area and are subjected to high levels of fishing mortality (Kramer & Chapman 1999). Mechanisms leading to adult export from MPAs are density-dependent movements of competitively subordinate individuals from preferred habitats (Dugan & Davis 1993, Abesamis & Russ 2005), seasonal migrations (McClanahan & Mangi 2000, Kaunda-Arara & Rose 2004) or random and home range movements or relocation (Rakitin & Kramer 1996, Kramer & Chapman 1999, Tupper 2007). Therefore spillover depends not only on biomass increase within protected areas but also on fish mobility and habitat characteristics around MPAs. Spatially, spillover should result in higher fish densities close to the borders of the MPA that decrease with distance, with specific patterns shaped by fish mobility and mortality imposed by fisheries in surrounding areas (Rakitin & Kramer 1996, Chapman & Kramer 1999, Russ et al. 2003). In fisheries adjacent to MPAs, spillover is expected to translate into aggregation of effort (Kellner et al. 2007) and increased catches (Murawski et al. 2005, Goñi et al. 2006, Abesamis et al. 2006a).

Theoretical studies of the effects of MPAs on fisheries (e.g. Gerber et al. 2003, Rodwell & Roberts 2004, Hilborn et al. 2006 and references therein) outnumber empirical evidence. These studies have generally assessed merits of MPAs for attaining single species yield stability and sustainability. In contrast the aims of empirical studies of MPA effects on fisheries have so far been modest, focusing on ascertaining the distribution of fishing effort and/or catches near MPAs. Several studies have demonstrated fishing effort aggregation along MPA boundaries (e.g. Yamasaki & Kuwahara 1989, McClanahan & Kaunda-Arara 1996, Johnson et al. 1999, McClanahan & Mangi 2000, Kelly et al. 2002, Willis et al. 2003, Beukers-Stewart et al. 2005, Goñi et al. 2006, Murawski et al. 2005), suggesting spillover of biomass to adjacent fisheries. To evaluate MPA effects on fishery catches some studies have monitored catch per unit effort (CPUE) over time in grounds adjacent to MPAs and have demonstrated increases in CPUE of target species after MPA creation (e.g. Yamasaki & Kuwahara 1989, McClanahan & Kaunda-Arara 1996, McClanahan & Mangi 2000, Roberts et al. 2001, Galal et al. 2002, Russ et al. 2004, Beukers-Stewart et al. 2005) or declines after MPA protection ended (Alcalá & Russ 1990). Other studies have measured gradients of CPUE across or near MPA boundaries, and found higher yields immediately adjacent to MPAs, with gradients extending from a few hundred to several thousand meters away for species of low to moderate mobility (e.g. McClanahan & Kaunda-Arara 1996, Rakitin & Kramer 1996, Russ & Alcalá 1996, Chapman & Kramer 1999, Kaunda-Arara & Rose 2004, Murawski et al. 2005, Abesamis et al. 2006a). Studies of MPA effects on lobster fisheries

have reported similar results (Kelly et al. 2002, Davidson et al. 2002, Goñi et al. 2006). With the exception of the lobster studies and those of Murawski et al. (2005) and Beukers-Stewart et al. (2005) which were conducted in temperate ecosystems, most of this work has been done on coral reef fisheries. In the Mediterranean Sea, only one study has evaluated spillover effects of an MPA on adjacent artisanal fisheries (Goñi et al. 2006), demonstrating effort concentration and gradients of lobster catches extending up to 1500 m from MPA boundaries.

MPA benefits to fisheries have been difficult to demonstrate empirically because studies are troubled by site specificity and lack of management replication and base-line data (Dugan & Davis 1993, Russ 2002). Also, few studies have investigated spillover for long enough periods to see the effects fully developed (Russ et al. 2004). Further, scales and effects of additional factors (e.g. habitat, depth) on density gradients confound interpretation (García-Charton & Pérez-Ruzafa 1999). As a result, no study has yet demonstrated net biomass export from an MPA to the adjacent fishery, although strong evidence is becoming available. For example Russ et al. (2004) demonstrated the build-up of reef fish biomass for over 2 decades adjacent to the Apo MPA in the Philippines, and Abesamis et al. (2006a) estimated the potential maximum contribution of spillover from the MPA at less than 10 % of the overall yield of the Apo Island fisheries. In a tag-recapture study Goñi et al. (2006) demonstrated the existence of negative gradients of CPUE of both tagged and untagged lobster from the boundary of a Mediterranean MPA where tagged lobster originated. More recently Tupper (2007) demonstrated net outward movement of tagged surgeonfish from an MPA in Micronesia, but little or no spillover of 4 other exploited fish species, and concluded that spillover was species-specific and influenced by reef topography.

Marine protected areas have become a popular management tool in the Mediterranean Sea, with the last census reporting 85 in the region (Ramos-Esplá et al. 2004). Research on the benefits of MPAs in the area has focused on rocky-littoral fish assemblages using underwater visual census (UVC) techniques, although some studies have targeted seagrass fish assemblages and invertebrate populations (Sánchez-Lizaso et al. 2000). The primary objective of these studies has been to evaluate population recoveries inside MPAs, and only recently has attention been directed to assessing density gradients across MPA boundaries. Biomass buildup inside Western Mediterranean MPAs has occurred, despite their relatively small size, because of previous high exploitation rates and the limited mobility of target species. Studies generally report significantly higher biomass inside than outside MPAs of Sparidae, Ser-

ranidae, Scorpaenidae, Labridae, Congridae, Muraeidae and Palinuridae species (Harmelin-Vivien et al. 1985, García-Rubies 1997, García-Charton et al. 2004, Jouvenel et al. 2004, Michelli et al. 2005, Planes 2005, Goñi et al. 2006, Ojeda-Martinez et al. 2007). Species from these taxa are either sedentary (Scorpaenidae, Muraenidae, Congridae) or exhibit low (Serranidae, Labridae) to moderate (Sparidae, Mullidae, Palinuridae) mobility (Harmelin 1987, Goñi & Latrouite 2005).

The objective of this study was to investigate spillover from 6 MPAs in the Western Mediterranean based on effects on artisanal fisheries. The study focused on fisheries operating fixed gears and targeting a variety of littoral species in and around 6 MPAs. Artisanal fisheries in the region are characterized by the use of particular gears targeting species or groups of species in certain habitats and seasons (i.e. 'fishing tactics'; see Pelletier & Ferraris 2000). Prior to this study, fleets and fishing tactics operating around the MPAs selected for study were largely unknown. Thus, our first objective was to characterize the fisheries associated with each MPA and to select fishing tactics targeting groups of species with different spillover potential. Then, we examined the spatial distribution of fishing effort and catch around the MPAs and hypothesized that, in the presence of biomass export, (1) fishing effort would concentrate on the boundary of the MPAs, and (2) catch per unit area would be highest near MPA boundaries and decrease with distance away from MPAs. We hypothesized that fisheries targeting Sparidae, Mullidae, Palinuridae, Serranidae, and Scorpaenidae species could benefit from spillover near MPAs (providing there was continuity of habitats through MPA boundaries), and that the spatial extent of spillover would vary from a maximum in fisheries catching species of moderate mobility (such as Sparidae) to a minimum in those catching species of very restricted mobility (such as Scorpaenidae).

MATERIALS AND METHODS

General approach. To test the hypothesis of biomass export we investigated the spatial distribution of effort density and catch per unit area as a function of distance from fishery closure (FC) boundaries. This approach follows that of several studies that have assessed spillover based on gradients of fish density and fishing effort away from MPA boundaries (e.g. McClanahan & Kaunda-Arara 1996, Russ et al. 2003, 2004, Murawski et al. 2005, Abesamis et al. 2006a). However, while those studies used CPUE as the density index, we used CUPA as a measure of production, as was also done by Goñi et al. (2006) and McClanahan & Mangi (2000), to consider potential local abundance

depletion that may be caused by aggregation of effort near MPA boundaries (Walters 2000). Depletion affects spatial CPUE patterns and can hamper assessing MPA effects to adjacent fisheries. We analyzed CUPA of the combined catch rather than the catch by species because artisanal fisheries in the region have nominal targets but aim at optimizing the aggregated catch.

This study was part of a multidisciplinary European research project (BIOMEX, <http://biomex.univ-perp.fr/>). In addition to the assessment of MPA effects on artisanal fisheries from 6 Western Mediterranean MPAs, the BIOMEX study assessed export of eggs/larvae and adult spillover through the evaluation of biomass gradients by UVC, baited video and experimental fishing (Planes 2005). Prior to this study, investigation of MPA fishery benefits in the Mediterranean had only been carried out on a single-MPA basis.

Study area. This study was carried out in the MPAs of Carry-le-Rouet Natural Park (hereafter called Carry), Cerbère-Banyuls Marine Natural Reserve (Banyuls), Medes Islands Marine Reserve (Medes), Tabarca Island Marine Reserve (Tabarca), Cabrera Archipelago National Park (Cabrera), and Cabo de Palos-Islas Hormigas Marine Reserve (Cabo de Palos) (Fig. 1). These MPAs were selected because they were all established at least 8 yr prior to our study (Table 1) and evidence of population recovery of exploited species was available (Carry: Harmelin et al. 1995; Banyuls: Bell 1983, Dufour et al. 1995; Medes: García-Rubies & Zabala 1990; Cabrera: García-Rubies 1997, Reñones et al. 1999; Tabarca: Bayle-Sempere 1999;

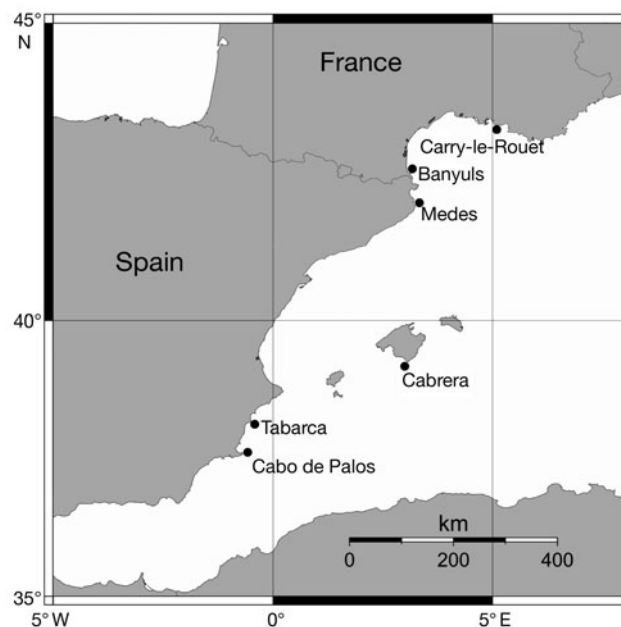


Fig. 1. Location of the western Mediterranean marine protected areas included in the study

Table 1. General characteristics of marine protected areas (MPA) selected for the study. Year = year of MPA establishment, FC = fisheries closure area, RU = restricted use area. Depth = maximum within areas. Main habitat types ranked based on proportion of area covered

MPA	Year	Type	Area (km ²)		Depth (m)		Main habitats	
			FC	RU	FC	RU	FC	RU & adjacent
Carry	1982	Coastal	0.85	0	30	30	Seagrass meadows Rocky photophylic	Seagrass meadows Rocky photophylic
Banyuls	1974	Coastal	0.65	5.85	45	60	Soft sandy Rocky photophylic Rocky coralligenous	Soft sandy Rocky photophylic Rocky coralligenous
Medes	1983	Island	0.93	4.18	60	60	Rocky coralligenous Rocky photophylic Soft sandy substrate	Soft sandy / muddy Seagrass meadows
Cabrera ^a	1991	Island	2.34	84.66	55	110	Rocky photophylic Seagrass meadows Soft sandy	Rocky photophylic Soft sandy Soft detritic / maërl
Tabarca	1986	Island	1.00	13.00	20	40	Seagrass meadows Rocky photophylic	Seagrass meadows Soft sandy Rocky photophylic
Cabo de Palos	1995	Coastal	2.70	16.28	80	100	Soft detritic / maërl Rocky coralligenous Rocky coralligenous	Soft detritic / maërl Rocky photophylic Rocky photophylic

^aDue to the large extension of the RU area of Cabrera, main habitats listed are RU habitats within 2 km of the FC boundaries

Cabo de Palos: García-Charton & Pérez-Ruzafa 1999, García-Charton et al. 2004). The selected MPAs are case studies for the Mediterranean, representative of islands or island groups and coastal strips protecting mainly shallow (<60 m) littoral ecosystems (Table 1). As in most Western Mediterranean MPAs, commercial fishery regulations include 2 zoning levels: (1) a fisheries closure (FC) area where all fishing activities are excluded, and (2) a restricted use (RU) area where some fishing is allowed, usually small scale artisanal fisheries (Table 1). Beyond these areas, MPAs are surrounded by open fishing waters. Carry does not have a RU area and the whole MPA is an FC. Similarly, in Tabarca fishing in the RU area is allowed only for pelagic species, thus for the purpose of this study the whole MPA is considered an FC. The size of FC areas among MPAs ranged from 0.65 to 2.7 km² and the size of RU areas from 0 to 84.66 km² (Table 1).

The spatial configuration and habitat characteristics vary among MPAs in the study. Cabrera presents sharper bathymetric gradients near FC boundaries than Medes and Cabo de Palos, and in Banyuls, Carry, and Tabarca gradients are minor (Fig. 2). Whereas in Cabrera the FC area consists of a string of unconnected bays, in the other MPAs it is a continuous area. The FC areas in Banyuls, Medes, Cabrera, and Cabo de Palos are all dominated by rocky habitats, while the RU and surrounding fishing areas are a mix of rock, gravel-sand, and sea-grass substrates, with the exception of Medes where RU and open fishing areas are dominated by soft bottoms (Table 1, Fig. 2). In Carry

and Tabarca FCs and surrounding areas *Posidonia oceanica* meadows are the dominant habitat (Table 1, Fig. 2). In all MPAs, habitats and depth in the RU areas are at least in some parts similar to those in the adjacent fishing areas (Fig. 2).

Sampling. Catch and effort data were collected during 2003 and 2004 onboard artisanal commercial fishing operations conducted within RU and open fishing areas adjacent to each MPA. For all fishing sets sampled, catch was identified to the species level and enumerated. Fish were measured and weighed. The date, geographic positions of the start and end of each fishing set, and occasionally intermediate positions, were recorded. For long-line fisheries the number and mean distance between hooks were recorded. For gill net and trammel net fisheries the length of the gear was recorded. During 2003, fisher interviews in nearby fishing harbors were also conducted. Since prior information on local fisheries was lacking, data collected onboard in 2003 covered all artisanal fishing activities in the study areas. These data, and information obtained during fisher interviews, were used to characterize fishing tactics operating in and around the MPAs and to select fishing tactics for data collection in 2004.

During 2003, a total of 212 fishing sets were sampled. Interviews and onboard sampling confirmed that fisheries around all MPAs were diverse and followed a variety of fishing tactics, and that the size of the vessels was small and fairly homogeneous (6 to 11 m). The most common tactics used gill nets, trammel nets, and long-lines. Between 5 and 14 fishing tactics were identified within

each MPA and nearby fishing grounds (49 in total). Based on the information gathered during 2003, we selected 14 fishing tactics on which to focus sampling during 2004 (Table 2). For the selection we used the following information and criteria: (1) the spatial dynamics of the fishing tactics, selecting those that occurred in grounds close to FCs, (2) the relative importance of each fishing tactic, selecting those with greatest fishing effort deployed by the largest number of boats, and (3) the species composition of catches, selecting those tactics catching highly exploited, non-migratory, benthic or necto-benthic species with suitable habitats within the FC areas. The selected fishing tactics used gill nets or a combination of gill nets and trammel nets to target various sparids, trammel nets to target mullids, scorpaenids, and palinurids, and bottom long-lines to target sparids and serranids (Table 2). Tactics targeting sparids were the most ubiquitous and used combined nets in Carry and Banyuls, gill nets in Medes and Tabarca, and long-lines in Tabarca and Cabrera (Table 2). The selected tactics operated both in RU and adjacent open fishing areas, ex-

cept in Carry and Tabarca where they only operated in open fishing areas and in Cabrera where they operated in the extensive RU area. Fishing tactics that were not selected for analysis targeted migratory species (e.g. Cabrera and Cabo de Palos: pelagic trammel nets) or species whose habitats were not present within FC areas (e.g. Carry and Banyuls: hake gill nets; Cabrera: lobster trammel nets). Other tactics were not considered because fishing effort was too low (e.g. Tabarca: sparid trammel nets; Cabrera: grouper hand line) or did not take place close to MPAs (e.g. Tabarca: cuttle fish trammel net; Cabo de Palos: dentex long-line).

During 2004, 488 fishing sets were sampled, of which we retained data from 399 for the spillover gradient analysis. Discarded sets were deployed too far from the potential areas of influence of the MPAs or were not directed to the fishing tactics selected for the study (boats could combine more than one fishing tactic on the same trip). Based on our estimates of total fishing days and number of boats operating around each MPA, our sampling effort covered about 5% of the

Table 2. Seasonality and catch composition of the fishing tactics selected for analysis of spillover in each marine protected area (MPA). 'Combined' tactics = gill and trammel net. Catch composition = proportion of retained catch (excluding pelagics), given for families that constituted $\geq 10\%$ by number. Target (*) and main non-target species are indicated in brackets

MPA	Fishing tactic	Season	Family	% total abundance
Carry	Sparids combined	Mar–Dec	Sparidae (<i>Sparus aurata</i> *, <i>Diplodus</i> spp., <i>Salpa salpa</i>)	68
	Mullet trammel net	Mar–Nov	Mullidae (<i>Mullus surmuletus</i> *)	33
			Scorpaenidae	16
			Labridae	10
Banyuls	Sparids combined	Apr–Sep	Sparidae (<i>Sparus aurata</i> *, <i>Pagellus</i> spp., <i>Diplodus</i> spp.)	44
	Lobster trammel net	Jun–Dec	Palinuridae (<i>Palinurus elephas</i> *)	63
			Uranoscopidae	10
Medes	Sparids gill net	Sep–Dec	Sparidae (<i>Pagellus erythrinus</i> *, <i>Pagellus bogaraveo</i> *)	41
	Mullet trammel net	May–Nov	Mullidae (<i>Mullus surmuletus</i> *)	29
			Sparidae	21
Cabrera	Grouper long-line	Sep–Dec	Serranidae (<i>Epinephelus marginatus</i> *, <i>Serranus cabrilla</i>)	23
			Sparidae (<i>Pagrus pagrus</i> *, <i>Dentex dentex</i>)	28
	Sparids long-line	Jan–May	Sparidae (<i>Pagrus pagrus</i> *, <i>Spondyllosoma cantharus</i>)	16
			Scyliorhinidae	58
	Rockfish trammel net	May–Aug	Scorpaenidae (<i>Scorpaena scrofa</i> *)	13
			Palinuridae (<i>Palinurus elephas</i> *)	10
Tabarca			Scyliorhinidae	12
			Sparidae	11
	Sparids long-line	Jan–Dec	Sparidae (<i>Diplodus</i> spp. *, <i>Pagrus pagrus</i> *)	33
			Moronidae	23
			Congridae	21
	Sparids gill net	Jan–Dec	Sparidae (<i>Dentex dentex</i> *, <i>Diplodus sargus</i> *, <i>Sparus aurata</i> *)	24
Cabo de Palos	Mullet trammel net	Jan–Dec	Mullidae (<i>Mullus surmuletus</i> *)	40
			Serranidae	11
			Sparidae	10
	Rockfish trammel net	May–Sep	Scorpaenidae (<i>Scorpaena scrofa</i> *, <i>S. porcus</i>)	37
			Sparidae	15
	Lobster trammel net	May–Sep	Palinuridae (<i>Palinurus elephas</i> *)	20
			Scorpaenidae	47
			Sparidae	19

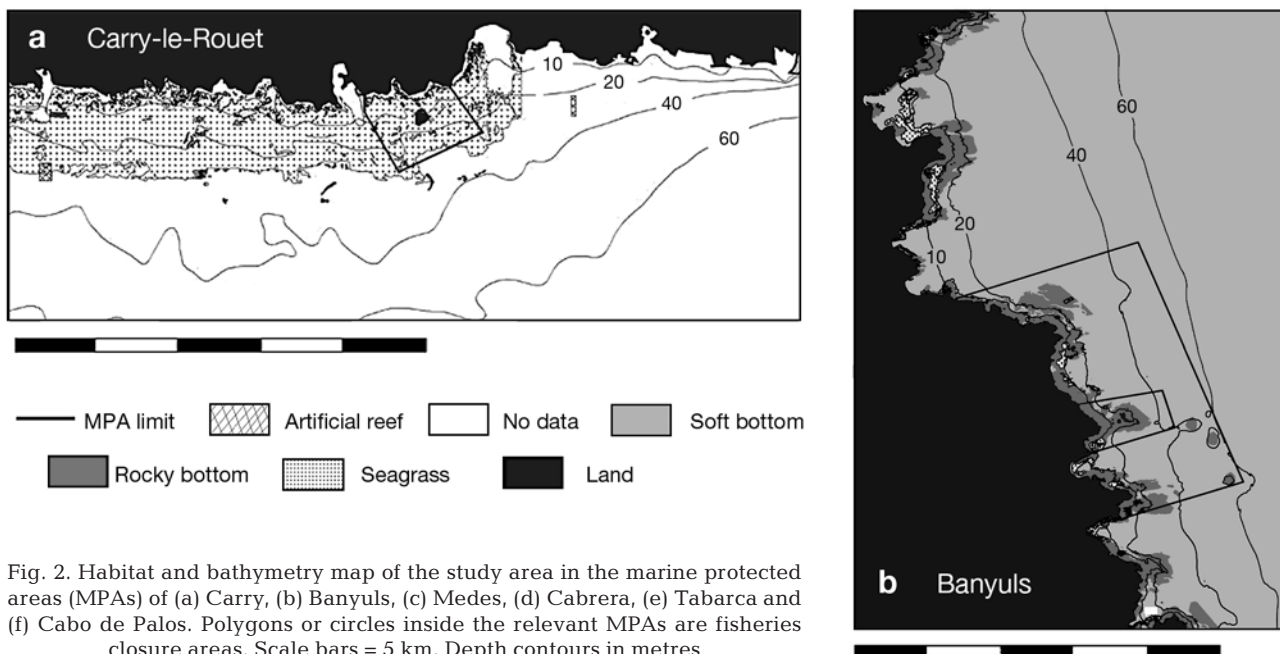


Fig. 2. Habitat and bathymetry map of the study area in the marine protected areas (MPAs) of (a) Carry, (b) Banyuls, (c) Medes, (d) Cabrera, (e) Tabarca and (f) Cabo de Palos. Polygons or circles inside the relevant MPAs are fisheries closure areas. Scale bars = 5 km. Depth contours in metres

total effort conducted by the selected fishing tactics in Carry, Tabarca and Cabo de Palos, 13% in Medes and Cabrera, and 33% in Banyuls.

The catches from the selected fishing tactics were highly diverse but dominated by a restricted number of species with moderate to low mobility. Discards were low and included low value species caught in small quantities or small pelagics caught accidentally. There were up to 76 species identified in a single tactic but 16 species belonging to 10 families made up 10% or more of the retained catch. The contribution of target taxa to the retained catch (by number, excluding pelagics) varied from 68% in the Carry sparid tactic to 13% in the Cabrera rockfish tactic (Table 2). Sparidae were the most abundant taxa in 11 of the 14 tactics studied.

Mean effort and retained catch per set ranged from generally lowest levels in long-line fishing tactics to highest among mullet trammel net fishing tactics. In the sparid and grouper long-line tactics, mean effort and retained catch ranged from 94 to 144 hooks and 4.1 to 10.5 fish per set, in the sparid gill net tactics from 627 to 1114 m of net and 5.7 to 17.3 fish per set, in the rockfish and lobster trammel net tactics from 591 to 1874 m of net and 3.5 to 50.6 individuals per set, and in the mullet trammel net tactics from 973 to 1578 m of net and 17.3 to 179.9 fish per set.

Data handling. The study area around each MPA was divided in square cells and for each fishing tactic we calculated effort density and catch per unit area (CPUA) by cell. We used GIS ESRI® ArcMap 8.3 and ESRI® ArcView 3.2 to map each study area, design a grid, and assign catch and effort data from the fishing

sets to the cells. Within each MPA, we defined the size of the cells based on an iterative process in search of a compromise between the length of fishing sets and the number of sets occurring within cells of alternative dimensions. Furthermore, the process was conducted under literature-informed constraint that the cell size should be likely to provide sufficient resolution to detect spillover. The length of fishing sets among MPAs and fishing tactics ranged from 100 to 5040 m with modes between 500 and 1500 m. We drew the set trajectories on the GIS maps based on start and ending positions and created polygons around each tracing. To consider the sinuosity of each set, we maintained the surface of each polygon proportional to the real length of the set (i.e. meters of gear deployed). The cell size selection process resulted in cells ranging from 150 × 150 m in Medes to 750 × 750 m in Cabo de Palos (Table 3). The total number of cells in which fishing sets occurred varied from 18 for Cabo de Palos lobster to 366 for Medes mullet tactics.

We assigned effort and catch from each fishing set to corresponding cells based on the overlay of polygons representing each set and the cells in each map, and allocated effort and catch per set to cells in proportion to the area of the corresponding polygon falling within each cell. For each fishing tactic we calculated effort density, CPUA and revenue by cell. Effort density was calculated as the sum of the number of hooks or meters of net, CPUA as the sum of the number of fish caught and retained, and revenue as the sum of the value of the retained catch in each cell. Mean effort density and CPUA per cell in the sparid and grouper long-line

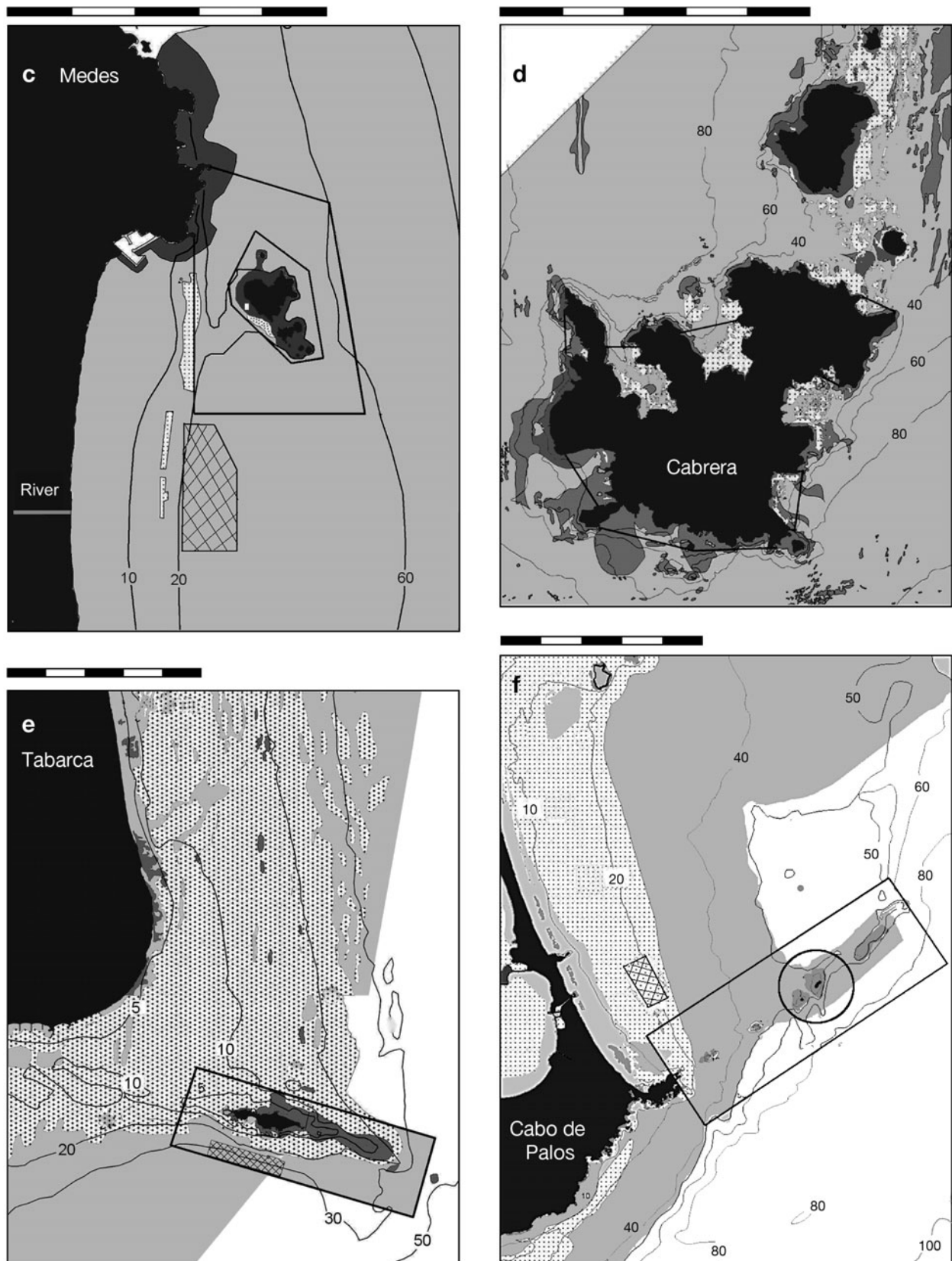


Fig. 2 (continued)

tactics ranged from 198 to 1065 hooks km⁻² and 15 to 88 fish km⁻², in the sparid gill net tactics from 1530 to 6006 m of net km⁻² and 39 to 962 fish km⁻², in the rockfish and lobster tactics from 1680 to 3595 m of net km⁻² and 7 to 105 individuals km⁻² and in the mullet tactics from 1602 to 5810 m of net km⁻² and 210 to 1112 fish km⁻² by cell.

The distance from the fished cells to the FC areas was measured by a straight line from the midpoint of each cell to the nearest FC boundary. For cells that crossed the FC boundary, distance was measured from the midpoint of the portion of the cell outside the FC. The maximum distance ranged from 1623 m for sparid and grouper tactics in Cabrera to 15690 m for sparid tactics in Tabarca. Depth was estimated from bathymetric maps and ranged from 5 m for the Carry mullet tactic to 100 m for the Cabrera sparid long-line tactic.

Data analysis. To investigate the spatial distribution of effort and catches we used a regression approach and explored the relationships of the following response variables: (1) effort density by cell (in meters of net or number of hooks per km²), (2) CUPA by cell (in number of fish caught and retained per km²), and (3) revenue by cell (in monetary value [€] of the CUPA per km²) with the explanatory variables (1) distance from the cell to the FC, and (2) mean depth of the cell. We introduced depth as a predictor variable to account for the potential effect of bathymetry on the distribution of species in the catch. First, as an exploratory tool, we used generalized additive models (GAMs, Hastie & Tibshirani 1990), as we expected relationships to be non-linear and this non-parametric technique allows

exploration of trends without constraining their shape. Distance and depth were introduced in the linear predictor as non-parametric smoothers with 4 degrees of freedom (df). For distances greater than 2000 m we added 1 degree of freedom every 1000 m. We determined appropriate probability distributions to implement the models by regressing the logarithm of the mean of each response variable in 200 m distance intervals, and the logarithm of the variance. For all fishing tactics and response variables the variance was closely proportional to the square of the mean; accordingly we used a gamma probability distribution. To relate the response variables to the linear predictor we selected the logarithmic link. To evaluate model fits we inspected the distribution of model deviance and residuals. Next, based on inspection and interpretation of GAM outputs for effort density and CUPA, we selected data for each fishing tactic within particular distances to FC boundaries and implemented generalized linear models (GLMs, McCullagh & Nelder 1989). Data selection was based on observing changes in slopes from trends away from FC boundaries; we used expert judgment rather than an objective criterion to define transition points. GLMs were implemented to estimate slopes and test the significance of linear relationships of effort density and CUPA with distance. The GLM linear predictors included distance to FC boundaries and depth. We used gamma and logarithmic link functions as for the GAM analyses. To evaluate the significance of each covariate, we performed analysis of deviance and tested each variable by comparing full models with models excluding the tested variable.

Within the analysis of deviance we tested the linearity of the depth covariate and introduced the term as a polynomial where appropriate. Tests were performed at the 95% confidence level. The GAM and GLM analyses were implemented using the gam(x) and glm(x) functions in S-Plus (Becker et al. 1988).

Table 3. Size of square cells in each marine protected area (MPA) and number of cells used in generalized linear models of effort density and catch per unit area with distance to fishery closures (FC) and depth by fishing tactic. Distance = distance from cell to nearest protection boundary (i.e. restricted use area boundary for Tabarca and FC boundary for other MPAs), depth = mean depth of cell, 'combined' tactics = gill net and trammel net

MPA	Cell side (m)	Fishing tactic	No. of cells	Distance range (m)	Depth range (m)
Carry	250	Sparids combined	34	40–995	5–50
		Mullet trammel net	45	5–998	5–50
Banyuls	300	Sparids combined	17	128–869	12–65
		Lobster trammel net	15	141–810	22–65
Medes	150	Sparids gill net	81	14–800	10–60
		Mullet trammel net	136	500–2497	10–35
Cabrera	250	Grouper long-line	91	20–1623	20–90
		Sparids long-line	38	712–1623	20–100
		Rockfish trammel net	66	40–1623	20–90
Tabarca	550	Sparids long-line	12	259–2490	15–60
		Sparids gill net	17	272–1922	10–30
		Mullet trammel net	18	270–3200	20–60
Cabo de Palos	750	Rockfish trammel net	21	229–1490	10–75
		Lobster trammel net	7	0–673	10–80

RESULTS

GAMs: Patterns of effort density, CUPA, and revenues adjacent to FC areas

Patterns of effort density and CUPA with distance to FCs were non linear with a general tendency to decline near boundaries (Figs. 3 to 8). The exception was the sparid long-line tactic in Cabrera, for which effort and CUPA were

initially stable and then declined (Fig. 6). The extent of effort density and CPUA gradients adjacent to FC areas among MPAs was 800 to 2490 m from FC boundaries for all sparid tactics, 998 to 3200 m for mullet tactics, 673 to 810 m for lobster tactics, and 1490 to 1623 m for rockfish and grouper tactics (Figs. 3 to 8). As distance from FC areas increased, effort and CPUA trends presented secondary peaks in the Carry (Fig. 3) and Medes (Fig. 5) sparid tactics and all the Banyuls (Fig. 4) and Tabarca (Fig. 7) tactics. Adjacent to FC boundaries, revenues followed the same trends as CPUA except for the Banyuls sparid (Fig. 4), Medes mullet (Fig. 5), Cabrera sparid and rockfish (Fig. 6) and Tabarca sparid long-line (Fig. 7) tactics. In these cases, revenues remained stable near boundaries while effort and CPUA declined. Based on the GAM patterns observed we selected data from cells within 673 to 3200 m from FC boundaries, depending on the MPA and fishing tactic, to run GLMs and estimate slopes of effort density and CPUA with distance to FCs (Table 3).

Effort, CPUA, and revenues were stable with depth within ranges fished by most fishing tactics (e.g. Fig. 3). Exceptions were the Carry sparid (Fig. 3a), Cabrera grouper and Cabo de Palos lobster tactics, in which effort, CPUA, and revenues peaked at intermediate depths, and the Cabrera rockfish tactic in which effort, CPUA and revenues were initially stable and declined in deeper waters (not shown).

GLMs: Gradients of effort density and CPUA as a function of distance to FC boundaries

Slopes of linear regressions of fishing effort density as a function of distance to FC boundaries for the 14 fishing tactics analyzed were negative and significant except for the Tabarca mullet trammel net tactic (Table 4). Slopes of regressions for CPUA with distance from the FC boundaries were also negative for the 14 fishing tactics analyzed and 9 were significant

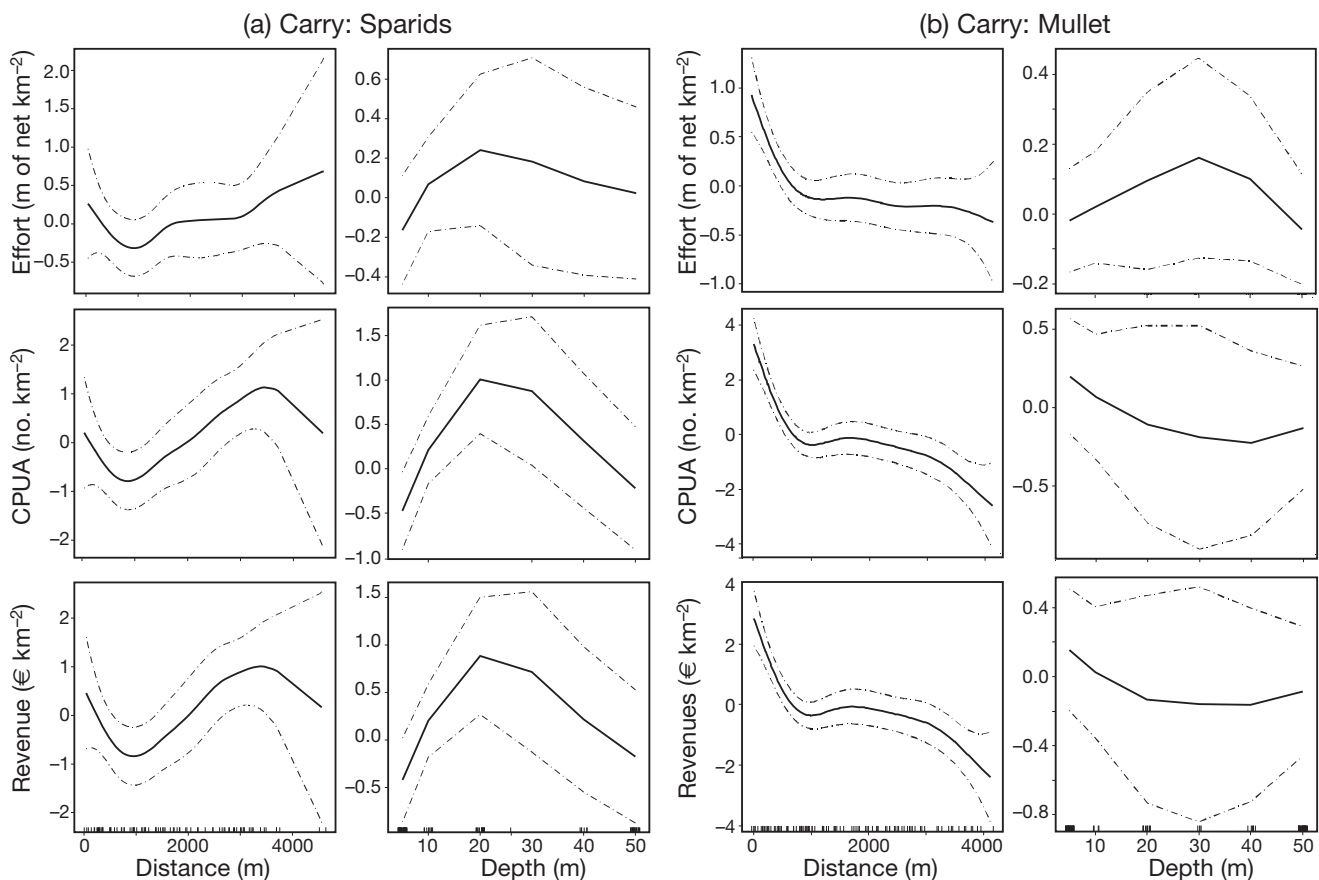


Fig. 3. Generalized additive model (GAM) predicted values by cell for effort (m of net km^{-2}), catch per unit area (CPUA, no. km^{-2}), and revenues (€ km^{-2}) within Carry-le-Rouet for (a) sparid and (b) mullet fishing tactics. Models incorporate the predictors 'distance to fisheries closure boundaries' and 'depth' as **nonparametric smoothers**, and **gamma variance and logarithmic link functions**. y-axes are standardized so that $y = 0$ represents the mean response. Bands around the predictions are approximate 95% confidence envelopes. Rug plot on the x-axis represents amount of data available

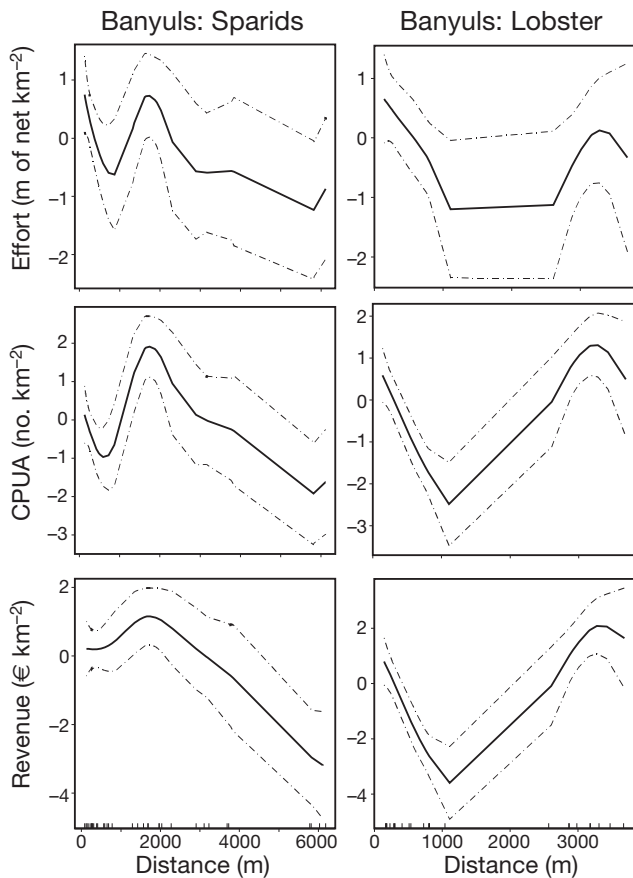


Fig. 4. GAM predicted values by cell for effort (m of net km⁻²), catch per unit area (CPUA, no. km⁻²), and revenues (€ km⁻²) within Banyuls sparid and lobster fishing tactics. Other details as in Fig. 3

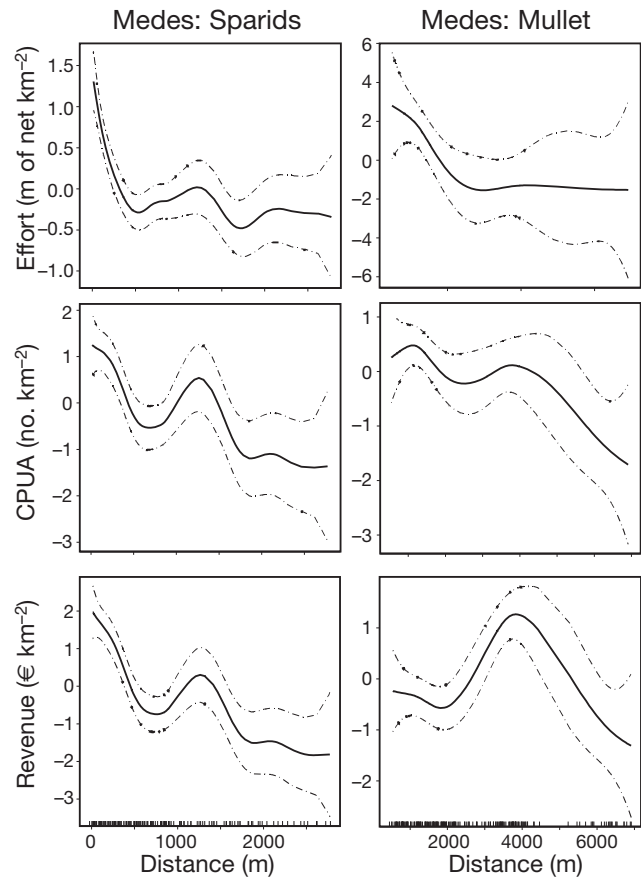


Fig. 5. GAM predicted values by cell for effort (m of net km⁻²), catch per unit area (CPUA, no. km⁻²), and revenues (€ km⁻²) within Medes sparid and mullet fishing tactics. Other details as in Fig. 3

(Table 4). Exceptions were the 3 Cabrera tactics ($p > 0.06$), the Tabarca mullet ($p > 0.1$) and the Cabo de Palos rockfish ($p > 0.4$) tactics (Table 4). Significant negative slopes of effort density and CPUA were steepest in the lobster tactic in Cabo de Palos and shallowest in the Cabrera rockfish and Medes mullet tactics (Table 4).

Effort density varied significantly with depth in the 3 fishing tactics in Cabrera and the rockfish tactic in Cabo de Palos ($p < 0.04$). CPUA varied significantly with depth in the grouper and rockfish tactics in Cabrera ($p < 0.04$). When significant, the relationship between effort density or CPUA and depth was non-linear, characterized by initial stable levels followed by a decrease, and best represented by a second degree polynomial. The effects of depth and distance were not correlated in the models for effort or CPUA, as the decrease with depth occurred after the decrease with distance was observed at several hundred meters away from FC boundaries.

DISCUSSION

In our study of 6 Western Mediterranean MPAs, concentration of fishing effort found near FC boundaries can be interpreted as spillover benefits to adjacent fisheries. Fishing effort was highest near the FC boundaries for all 14 fishing tactics in the 6 MPAs studied and declined significantly in all but the mullet trammel net tactic in Tabarca. Where observed, this pattern of 'edge fishing' or 'fishing the line' around MPAs has been taken as a clear indication of spillover (e.g. Yamasaki & Kuwahara 1989, McClanahan & Mangi 2000, Kelly et al. 2002, Willis et al. 2003, Murawski et al. 2005, Goñi et al. 2006), although effort aggregation can also result from redistribution of effort after MPA creation (Halpern et al. 2004, Murawski et al. 2005), or from perceived benefits leading fishers to occupy fishing spots along boundaries even when yields become locally depleted (McClanahan & Kaunda-Arara 1996, McClanahan & Mangi 2000, Willis et al. 2003). In the case of well established MPAs such as

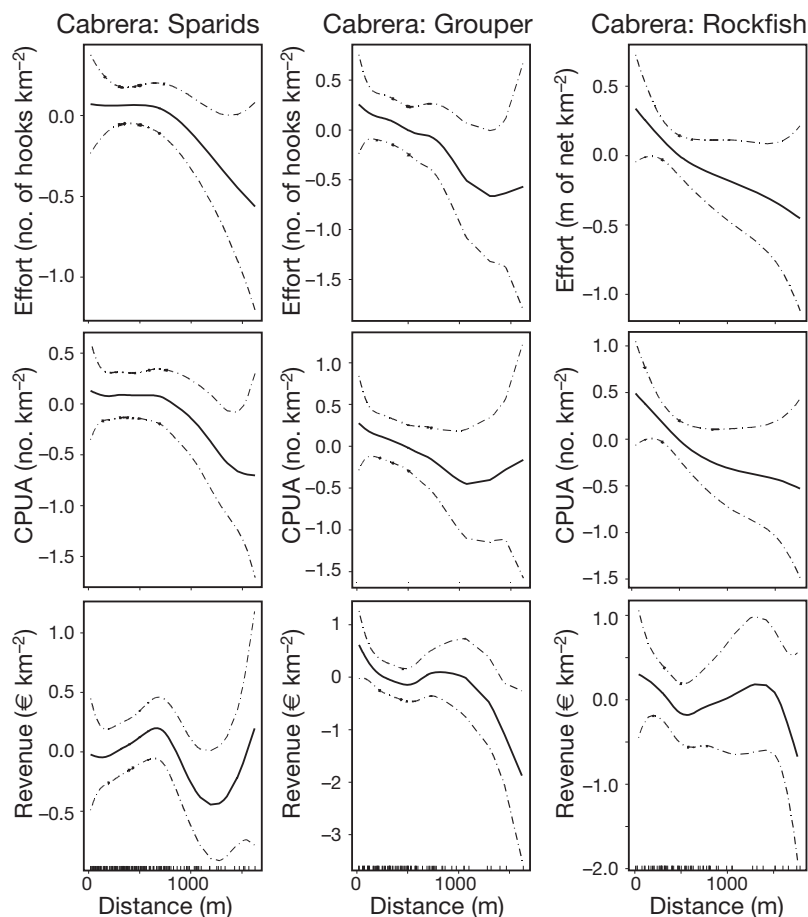


Fig. 6. GAM predicted values by cell for effort (no. of hooks or m of net km^{-2}), catch per unit area (CPUA, no. km^{-2}), and revenues (€ km^{-2}) within Cabrera spard long-line, grouper long-line, and rockfish trammel net tactics. Other details as in Fig. 3

those in this study we can safely assume that observed effort distribution responded to achieved fishery benefits, as predicted by MPA fishery models (Maury & Gascuel 1999, Kellner et al. 2007). Aggregation of effort near MPA boundaries is not always observed when effective protection takes place. Such cases have been reported when the cost from traveling from home ports to MPAs is high (Wilcox & Pomeroy 2003), or when greater yield variability near MPAs attenuates spillover benefits (Abesamis et al. 2006a). Furthermore, MPA boundaries may not attract fishers when there are productive fishing grounds nearby (Abesamis et al. 2006a). In our study, effort aggregation near FC boundaries occurred even when productive fishing grounds were found in the vicinity of several MPAs, as revealed by secondary effort density peaks in GAM plots. Only for the mullet trammel net tactic in Tabarca, important fishing grounds located about 7 km north of the MPA (Figs. 1 & 7) may have detracted effort from the MPA boundaries.

Trends in fishery production in this study declined with distance from FC boundaries and are generally consistent with reported build-up of exploited species in the study MPAs. CPUA was

Table 4. Slopes of effort density and retained catch by area (CPUA) as a function of distance to fishing closure (FC) boundaries from generalized linear models incorporating distance and depth. Distance was to restricted use area boundary for Tabarca and to FC boundary for other MPAs. Models included a gamma distribution and a logarithmic link. 95% CI are confidence intervals. *indicates significant slopes at the 95% level. Combined tactics are gill net and trammel net

MPA	Fishing tactic	Effort density			CPUA		
		Slope	95%CI	Prob. of <i>F</i>	Slope	95%CI	Prob. of <i>F</i>
Carry	Sparids combined	-0.0011	-0.002, -0.0002	0.0076*	-0.0018	-0.0034, -0.0002	0.0221*
	Mullet trammel net	-0.0012	-0.002, -0.0004	0.0019*	-0.0052	-0.0069, -0.0035	0.0001*
Banyuls	Sparids combined	-0.0034	-0.0055, -0.0013	0.0012*	-0.0027	-0.0052, -0.0002	0.0238*
	lobster trammel net	-0.0014	-0.0027, -0.0001	0.0503*	-0.0037	-0.0059, -0.0015	0.0001*
Medes	Sparids gill net	-0.0018	-0.0026, -0.001	0.0001*	-0.0026	-0.0037, -0.0015	0.0001*
	Mullet trammel net	-0.0005	-0.0007, -0.0003	0.0009*	-0.0007	-0.0013, -0.0001	0.0399*
Cabrera	Grouper long-line	-0.0006	-0.0009, -0.0003	0.0089*	-0.0004	-0.0008, 0.0000	0.0635
	Sparids long-line	-0.0017	-0.0033, -0.0001	0.0059*	-0.0012	-0.0027, 0.0003	0.0697
	Rockfish trammel net	-0.0004	-0.0007, -0.0001	0.0329*	-0.0004	-0.0009, 0.0001	0.1172
Tabarca	Sparids long-line	-0.0017	-0.003, -0.0004	0.0114*	-0.0014	-0.002, -0.0008	0.0015*
	Sparids gill net	-0.0011	-0.0015, -0.0007	0.0123*	-0.0008	-0.0001, -0.0015	0.0451*
	Mullet trammel net	-0.0002	-0.0006, 0.0002	0.1849	-0.0002	-0.0006, 0.0002	0.1561
Cabo de Palos	Rockfish trammel net	-0.0011	-0.0017, -0.0005	0.0036*	-0.0001	-0.0018, 0.0017	0.4881
	Lobster trammel net	-0.0071	-0.017, -0.0035	0.0027*	-0.0077	-0.0118, -0.0036	0.0017*

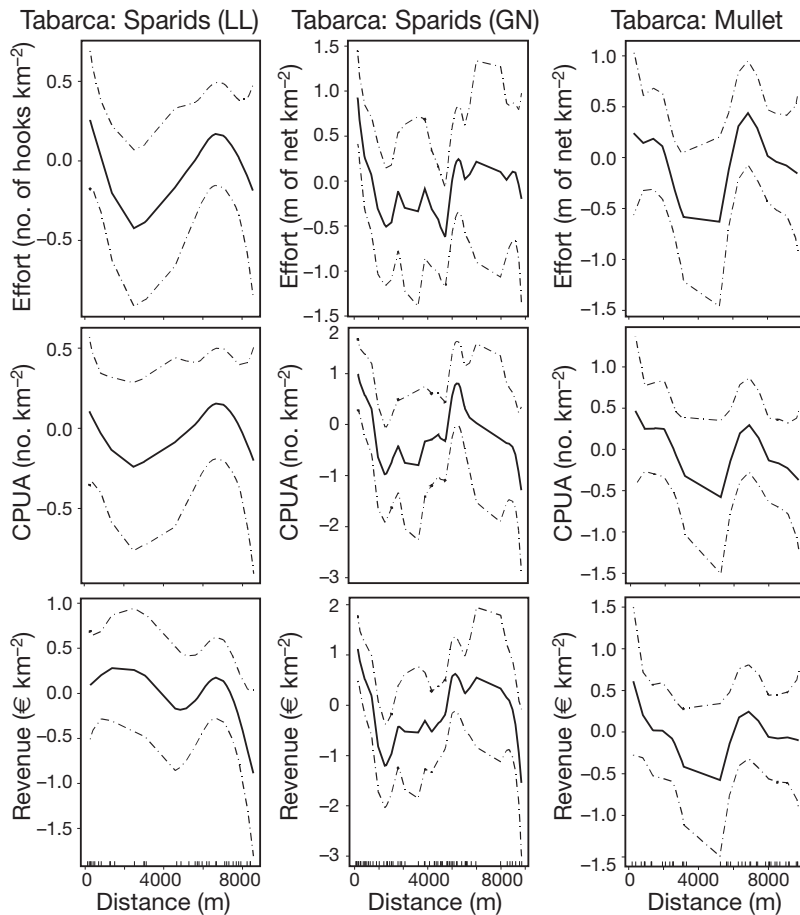


Fig. 7. GAM predicted values by cell for effort (no. of hooks or m of net km^{-2}), catch per unit area (CPUA, no. km^{-2}), and revenues (€ km^{-2}) within Tabarca sparid long-line (LL) and gillnet (GN) and mullet fishing tactics. Other details as in Fig. 3

highest near FC boundaries for all 14 fishing tactics, and in 9 of the cases negative slopes away from boundaries were significant. Exceptions were the 3 tactics in Cabrera, and the Tabarca mullet and Cabo de Palos rockfish tactics. Build-up of exploited species in MPAs in this study is well documented (e.g. García-Rubies 1997, Sánchez-Lizaso et al. 2000, García-Charton et al. 2004, Ojeda-Martínez et al. 2007) and contemporary underwater visual census (UVC) of fish density indicated declining gradients from inside FCs to the adjacent fishing areas of sparids (*Diplodus* spp., *Sparus aurata*, *Dentex dentex*, *Pagrus pagrus*, *Spondylus cantharus*), serranids (*Epinephelus* spp., *Serranus* spp.), labrids (*Labrus* spp., *Symphodus* spp.), scorpaenids (*Scorpaena scrofa*), muraenids (*Muraena helena*), and congrid (*Conger conger*) (Planes 2005). These are the main species caught by fishing tactics in this study and thus, our findings support the biomass export hypotheses in particular for the sparids tactics (Table 2). Other species were not assessed by

UVC due to their behavior (e.g. *Pagellus* spp., *Scyliorhinus canicula*) or cryptic nature (e.g. *Scorpaena porcus*). In tactics where these species are important, in particular in the Medes sparid tactic, the lack of UVC evidence makes it difficult to assess how production gradients relate to population build-up. Evidence from UVC on species caught in all 3 tactics in Cabrera indicates density build-up in the FCs and thus biomass export could have been affected by habitat characteristics or by methodological factors also related to habitat. The only species reported to be more abundant outside than within FC boundaries was *Mullus surmuletus* in Medes (Planes 2005), where the species is targeted by the trammel net fishing tactic.

Fishery production gradients in our study were generally consistent with expected MPA benefits based on behaviour of species in the catch and fishery exploitation. Spillover benefits are expected from MPAs for species that exhibit moderate movements and that are highly exploited when species are protected in habitats with continuity in adjacent areas (Kramer & Chapman 1999, Tewfik & Bene 2003, Tupper 2007).

Habitat continuity should not be an obstacle for biomass export in our study as we selected tactics among MPAs that target species for which suitable surroundings were present to varying degrees in all the FCs and adjacent fished areas. These habitats are coastal rocky and mixed substrates for sparids, scorpaenids, labrids and serranids (Harmelin 1987) present in all MPAs, *Posidonia* meadows for *Mullus surmuletus* (Harmelin 1987) abundant in Carry and Tabarca and rocky and coralligenous habitats for *Palinurus elephas* and associated species (Goñi & Latrouite 2005) present in Banyuls and Cabo de Palos. Trends were generally significant among fishing tactics catching predominantly moderately vagile sparids, mullids, and palinurids, while trends were not significant in the tactics with presence of varying proportions of sedentary species, in particular scorpaenids (Table 2). Thus, significant production trends in sparids, mullet and lobster tactics are a likely indication of spillover benefits, while non-significant negative trends in rockfish tactics may suggest lesser benefits to the fisheries. The influence of fishery exploitation was not assessed in our study, but production gradients were consistent with gear efficiency. Theory predicts that when populations

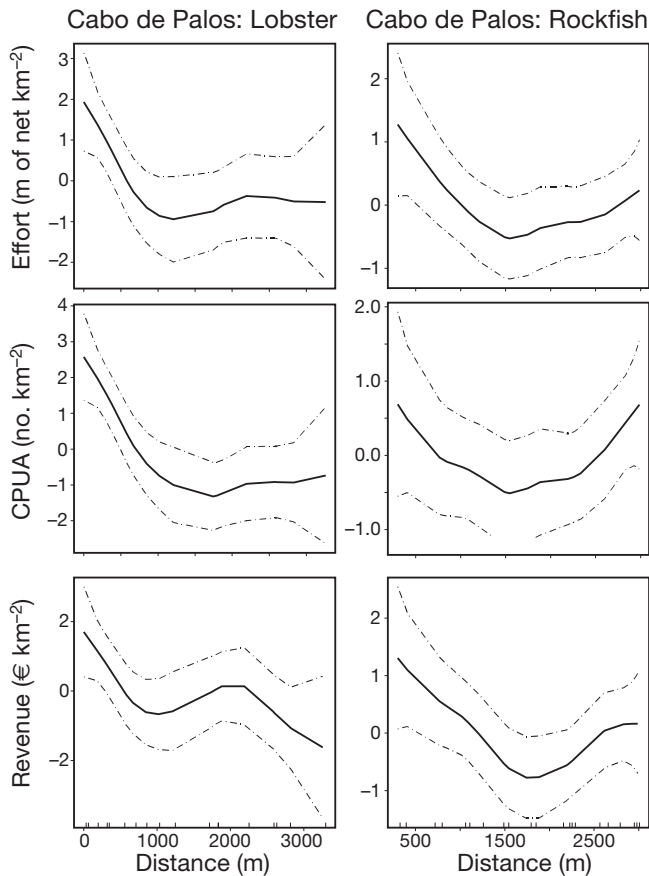


Fig. 8. GAM predicted values by cell for effort (m of net km^{-2}), catch per unit area (CUPA, no. km^{-2}), and revenues (€ km^{-2}) within Cabo de Palos rockfish and lobster fishing tactics. Other details as in Fig. 3

build up inside MPAs, density gradients away from FCs are, inter alia, a function of species catchability (Rakitin & Kramer 1996), and magnitude of spillover effects observed in tropical MPAs have been partially attributed to the differential efficiency of fishing gears used in adjacent fisheries (McClanahan & Mangi 2000, Abesamis et al. 2006a). In our study the Carry mullet tactic exhibited one of the steepest gradients and used trammel nets, the most efficient gear in the study. The steep production trends in the lobster tactics are also consistent with the high catchability of the target species in trammel nets (Goñi et al. 2003). On the other hand, the sparid tactics exhibited shallower production gradients, congruent with lower efficiency of gill nets and long-lines that allow fish to move greater distances before being caught.

Production gradients of some tactics in this study were probably affected by characteristics of habitats within and around MPAs. Gradients in CUPA were not significant for the 3 tactics in Cabrera, and also for the Tabarca mullet and Cabo de Palos rockfish tactics, despite reported biomass build-up of the most common

species in the FCs. In Cabrera, habitat characteristics pose challenges for the evaluation of benefits to fisheries based on gradients due to the spatial configuration of the FC areas, which consist of numerous bays around a main island. Regardless of general availability of suitable areas within and around FCs, the relatively secluded bays may conceal features not present in adjacent areas that may affect biomass export of some species. Further, since it was not possible to identify specific bays where individuals originated, distances measured from fishing set locations to the nearest bay might not accurately represent the distance between the protection boundary and the fishing location. As with effort, production gradients in the Tabarca mullet tactic may have been affected by the presence of productive fishing grounds 7 km north of the MPA. In Medes, the opposite was observed, as gradients of effort and production of the mullet tactic tested significant despite reported absence of biomass build-up of *Mullus surmuletus*. In this case gradients were affected by the presence of *Posidonia* meadows near the MPA southeast boundary which were scarce within the FC area. Thus, we do not interpret these gradients as indicative of spillover because of the reported lack of build-up and the effects of habitat on gradients.

The spatial extent of production gradients away from FC boundaries in our study is consistent with spillover effects reported for littoral and coral reef species in the Mediterranean and are low compared to reports for soft bottom temperate species. Effects ranged from about 700 to 1000 m in the lobster and mullet tactics, up to 2500 m in the sparid tactics. For the lobster tactics, the scale was similar to that for *Palinurus elephas* (1000 to 1500 m; Goñi et al. 2006) from a western Mediterranean MPA. Fish density gradients from UVC observations in the MPAs in this study extended only up to 500 m from the boundaries of FCs (Planes 2005), but UVC likely underestimates the potential significance of fish exported to the local fisheries since only fish in excess of those caught are detected (Russ & Alcala 1996). Gradients of CPUE away from MPAs similar to those reported in our study have been documented for related species in coral reefs (e.g. McClanahan & Mangi 2000, Roberts et al. 2001, Galal et al. 2002, Russ et al. 2004, Abesamis & Russ 2005, Abesamis et al. 2006b). More extensive CPUE gradients extending up to 4 km from MPA boundaries have been reported in the Northwest Atlantic for soft bottom groundfish species (Murawski et al. 2005), and up to 13 km in waters off Japan for the crab *Chionoecetes opilio* (Yamasaki & Kuwahara 1989).

Fisheries revenues generally declined with distance to fisheries closures in a similar way to production but cannot be easily compared with the few studies avail-

able, which have reported varying results. In our study, enhanced revenue gradients were expected given the greater size (and value) of many exploited species near FCs (Planes 2005). The exception was in Cabrera, where the mean size of species important in the catch of the 3 fishing tactics (sparids, e.g. *Pagrus pagrus*, *Spondyllosoma cantharus*; scorpaenids, e.g. *Scorpaena scrofa*) were smaller near protection boundaries due to the presence of nursery habitats in the FC areas (Planes 2005). Revenue gradients were also influenced by variation in the species composition of catches with distance to FC boundaries, as occurred in the Banyuls and Tabarca sparid tactics. In previous studies, Murawski et al. (2005) reported higher economic yields near temperate MPAs while McClanahan & Kaunda-Arara (1996) observed the opposite in an MPA in Kenya; interestingly effort concentrated along boundaries in both MPAs. Kelly et al. (2002) found that economic yields of lobster near an MPA in New Zealand were similar to those in unprotected reefs (despite the lesser quality of habitat and concentration of effort near the MPA) and interpreted these results as benefits to the fishery. Finally, Abesamis et al. (2006a) reported higher but very variable economic yields near protection boundaries in the Apo reserve in the Philippines, which caused effort not to concentrate there. Thus, gradients in revenues can be independent of effort trends and may reflect particular conditions within and around MPAs.

Including non-linear terms in our analysis was instrumental to elucidate trends of fishing effort and production. In a similar study, and to avoid the confounding influence of neighboring MPAs on fishery gradients, Murawski et al. (2005) selected distances for the study upon visual examination of trends of catch and effort data. In this study GAMs helped to identify the range at which protection effects operated and recognize habitat discontinuities or 'hot spots' (areas of high production caused by particularly beneficial habitat conditions) located in the vicinity of the MPAs. Secondary peaks of effort and production away from FC boundaries were detected for several tactics. These peaks were in some cases associated with the presence of traditional fishing grounds around the MPAs. In the sparid tactics of Carry and Banyuls (Figs. 3b & 4), these grounds corresponded with particular capes, and in the lobster tactic in Banyuls (Fig. 4) with an isolated coralligenous zone within extensive sandy bottoms. For Medes mullet (Fig. 5) and Cabo de Palos rockfish (Fig. 8) tactics, secondary peaks coincided with the presence of artificial reefs adjacent to the RU boundaries (Fig. 2). In Tabarca, secondary peaks at about 7 km from the MPA (Fig. 7) were associated with traditional fishing grounds around particularly heterogeneous habitats. If not detected, secondary peaks could

have obscured the analysis of effort and production trends in relation to FCs.

Recreational activities around MPAs may have weakened effort and production gradients in artisanal fisheries found in our study but would not invalidate our results. Recreational fisheries are likely to compete with artisanal fisheries for biomass export benefits and might level trends of effort and production away from FCs. Hook and line recreational fisheries could have affected catches in this study because the target species are also caught by most fishing tactics studied (e.g. *Serranus* spp., *Diplodus annularis*, *D. sargus*, *D. vulgaris*, *Sparus aurata*, *Spondyllosoma cantharus*) (Harmelin et al. 1995). Also spear-fishing, very popular in the western Mediterranean (Coll et al. 2004), might compete with the sparid, grouper and rockfish tactics. Further, non-consumptive uses such as recreational diving might create conflicts and affect choice of fishing location within RU areas. In Cabo de Palos, for example, fishermen prefer to fish in the rocky shoals least frequently visited by divers (S. Polti, University of Murcia, pers. comm.). Assessing the effects of recreational activities on gradients of artisanal fisheries was beyond the scope of this study, but they should be considered in further studies of western Mediterranean MPAs. In our study, recreational fisheries should not invalidate conclusions of spillover but could have lessened the magnitude of the observed effects.

This study highlights the complexities of analyzing biomass export from artisanal fisheries but benefits from a design that included multiple MPAs and fisheries. Studies of MPA effects are often limited by the lack of baseline data and management replication, but lack of replication was alleviated here by the implementation of the same sampling design in 6 MPAs within the same region. Despite differences among the MPAs studied here, their resemblances are underscored by the similarity of the fisheries operating around them. We conclude that concentration of fishing effort and significant negative gradients of fishery production from artisanal fisheries in the study can be best explained by processes of spillover. Our interpretation is backed up by UVC studies in the same MPAs which reported higher biomass of the main species caught in fisheries inside MPAs than in fished areas in all 6 MPAs studied (Planes 2005). The spatial extent of spillover effects was consistent with species mobility and fisheries efficiency. Production gradients were not significant when characteristics of habitat or environmental conditions within and around MPAs imposed challenges for trend evaluation. These conclusions are strengthened by the composite findings of the several cases studied.

Our results suggest that coastal MPAs can be an effective management tool for artisanal fisheries in the

region and can be extended to the rest of the western Mediterranean, as the fishing tactics studied are typical of the region and represent an important component of the total fishing activities (Alarcón 2001, Colloca et al. 2004). Consistent with conditions identified for MPAs to benefit fishing yields (Hilborn et. al 2004), these fisheries target multispecies complexes dominated by species of restricted mobility. This study is a step forward in evaluating the effects of fishing closures in Mediterranean fisheries.

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