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# EVALUATION OF THE RESERVE EFFECT IN A MARINE PROTECTED AREA IN SAGRES (PNSACV) 

Mestrado em Biologia Marinha<br>Trabalho efetuado sob a orientação de:<br>Barbara Horta e Costa (CCMAR, UAlg) e Prof. Jorge Gonçalves (CCMAR, UAlg)

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#### Abstract

Marine Protected Areas (MPAs), especially when set up as marine reserves, have been found beneficial for fish communities and species worldwide. Evaluation of MPA effects needs to be done to understand whether existing protection measures are efficient or not. To detect potential protection effect, we aimed to assess differences in demersal fish and commercial invertebrate community at Ilhotes do Martinhal marine reserve. Based on a comparison between locations situated inside and outside the reserve, differences in richness, abundance, length and biomass were analysed at community and species level. The influence of physical habitat was investigated, as it could get confounded with protection effect. In addition, we wanted to validate the results from Stereo Baited Remote Underwater Video (SBRUV) by another method, being Stereo Diver Operated Video (SDOV). Comparison between methods performance and costs was held to decide on the more efficient monitoring tool. SBRUV results suggest that the marine reserve provides positive effects, especially as it sustains greater biomass of target species and target species above minimum landing size. The reserve was found beneficial for Diplodus sargus, a valuable commercial, and Labrus bergylta, a by-catch species. However, two species showed an opposite pattern, indicating negative protection effect or influence of other habitat characteristics rather than physical complexity. Furthermore, SBRUV results for abundance were non-conclusive of differences, probably due to a delayed response. SDOV showed no signs of positive protection effects, with some results complementary while other contradictory to SBRUV's ones. This was attributed to the differences in community sampled, reflecting diver and bait effects. As illegal fishing gear was encountered inside the reserve, legal enforcement and active management might play a key role in future reserve success. This study is especially relevant for further monitoring and revaluation of protection measures and zonation of Ilhotes de Martinhal.


Keywords: marine protected areas, reserve effect, stereo baited cameras, diver-operated video, demersal fish assemblages

## Sumário Executivo

As Áreas Marinhas Protegidas (AMPs) são consideradas uma ferramenta de conservação de meio marinho que tem como objetivo principal a proteção de habitats, de ecossistemas e da biodiversidade. Além disso, as AMPs podem promover a pesca sustentável e a recuperação de recursos de pesca. Especialmente quando designadas como reservas marinhas, onde todas as atividades extrativas são proibidas, as AMPs tem sido benéficas para comunidades e espécies de peixes. Os efeitos positivos geralmente associados a AMPs são o aumento de riqueza de espécies, de densidades, de tamanhos e de biomassa de peixes e invertebrados. Para avaliar os efeitos de medidas de proteção, idealmente, são realizadas comparações de dados de antes e de depois da sua implementação. Na ausência de dados anteriores à implementação de uma AMP, uma comparação entre locais com proteção (impacto) e sem proteção (controle) pode usar-se, tendo em conta a influência de habitat que pode confundir-se com efeitos de proteção. Neste estudo, foi comparada a comunidade de peixes demersais e invertebrados comerciais dentro e fora de uma reserva marinha dos Ilhotes do Martinhal em Sagres. Esta zona foi estabelecida em 2011 como área de proteção parcial I (a única atividade extrativa permitida é a apanha de percebe nas arribas da costa por pescadores comerciais, sendo considerada reserva marinha para o subtidal) integrante no Parque Marinho do Parque Natural do Sudoeste Alentejano e Costa Vicentina (PNSACV). Como os dados comparáveis anteriores à implementação da reserva não estavam disponíveis, os dados sobre a comunidade de peixes demersais e invertebrados comerciais foram comparados entre locais da reserva e locais adjacentes (zona complementar ou 'buffer') com substrato rochoso. A influência da complexidade física de habitat rochoso foi também considerada na análise porque os efeitos de reserva podem confundir-se com efeitos de habitat. As diferenças na riqueza de espécies foram investigadas. Em relação à abundância e biomassa, as diferenças foram analisadas por grupos de espécies (espécies demersais, espécies comerciais, espécies comerciais acima de tamanho mínimo legal (TML) e abaixo de TML, espécies sem interesse comercial). Ao lado da abundança e da biomassa, as diferenças de tamanhos foram também examinadas nas espécies individuais mais abundantes: comerciais (Diplodus sargus e Diplodus vulgaris) e sem interesse comercial (Coris julis, Serranus cabrilla, Labrus bergylta).

Dois métodos estéreo-vídeo de recolha de dados foram usados: as Câmaras Iscadas Estéreo (SBRUV) e o Vídeo Estéreo por Mergulho (SDOV). Estes métodos apresentam uma alternativa ao Censo Visual por Mergulho (UVC) que é o método o mais usado em monitorização de AMPs. Os dois métodos estéreo-vídeo também são de caracter não destrutivo, e por isso podem
ser preferíveis a uma monitorização de reservas marinhas quando comparados com as artes de pesca. Os dois métodos permitem estimar tamanhos de organismos mais precisos que os estimados por observações directas em UVC e também oferecem uma gravação permanente que pode ser vista por múltiplos observadores no laboratório. Assim, o erro ligado ao observador é reduzido quando comparado ao UVC. Dado a presença de isco, as SBRUVs servem como ponto de atração para peixes e como resultado, este método é considerado eficiente para a amostragem de riqueza de espécies de peixes, e particularmente de peixes carnívoros de tamanhos grandes.

Os resultados de amostragem com SBRUV sugerem que existem efeitos positivos dentro da reserva, visto ter-se encontrado uma maior riqueza e uma maior biomassa da comunidade de peixes demersais e de invertebrados comerciais dentro da reserva em relação à zona adjacente. Principalmente, a biomassa de espécies comerciais e de espécies comerciais acima de TML, os dois grupos que se espera que respondam mais à exclusão de pesca, foi mais alta dentro que fora da reserva. Os resultados sugerem que a espécie $D$. sargus, uma das principais espécies alvo nesta região, terá beneficiado da proteção em termos de tamanhos e de biomassa. A abundância e a biomassa de L. bergylta, que é uma de espécie de tamanho grande, mas que é capturada de forma acessória, também foram significativamente maiores dentro da reserva em comparação com zona adjacente. No entanto, Diplodus vulgaris (também de interesse comercial) e $S$. cabrilla (sem ou de baixo interesse comercial) parecem mais associados aos locais fora da reserva, sugerindo uma resposta negativa à proteção. Os resultados negativos nestas espécies podem também estar ligados a características do habitat sem ser a complexidade física. Neste estudo, não se encontraram diferenças significativas entre a complexidade física de habitats dentro e fora da reserva, nem interações significativas entre proteção e habitat. No entanto, algumas espécies apresentaram uma maior afinidade aos habitats de menor (D. sargus) ou de maior complexidade física ( $D$. vulgaris abaixo de TML).

Apesar dos resultados significativos para a biomassa, os resultados da abundância de peixes e invertebrados não revelaram diferenças suficientes para apoiar os efeitos positivos de proteção. No geral, os tamanhos e a biomassa fornecem uma resposta mais rápida relativamente as medidas de proteção, resultando diretamente da eliminação da pesca que selecciona os indivíduos de tamanhos maiores. Os efeitos em abundância podem precisar de mais anos de proteção para serem reconhecidos, devido à variabilidade de fatores ambientais que afetam o sucesso de recrutamento em organismos marinhos. Os resultados da abundância podem também ter uma ligação com a pesca ilegal, visto que artes de pesca ilegais (covos) foram encontradas repetidamente dentro da reserva. Neste contexto, a implementação de uma gestão ativa, baseada
na participação da comunidade local, na fiscalização e na monitorização ao prazo longo será fundamental para segurar o sucesso da reserva no futuro.

Em relação à comparação entre os dois métodos, o SBRUV mostrou-se eficiente para monitorização, visto que este método permitiu a deteção de eventuais efeitos de reserva em riqueza e em biomassa. Os resultados de SDOV não mostraram diferenças entre níveis de proteção que sugerissem efeitos positivos de reserva. Contudo, os resultados do SDOV validaram os resultados de SBRUV ao terem também detetado que os adultos de $D$. vulgaris preferiam os locais da parte oeste fora da reserva em comparação com os locais dentro da reserva. Ao contrário e como resultado complementar ao SBRUV, o SDOV revelou que, na reserva, foi observada uma maior proporção de juvenis desta espécie que nos locais adjacentes. Estes resultados sugerem que os indivíduos de $D$. vulgaris partem da reserva quando adultos. Relativamente a comunidade amostrada, o SDOV amostrou as espécies mais comuns e conspícuas (por exemplo $D$. vulgaris) com sucesso, mas já não foi eficaz na amostragem de espécies mais crípticas (por exemplo Symphodus spp. ou S. cabrilla). No total, o SDOV forneceu uma riqueza de espécies menor quando comparado ao SBRUV. Estes resultados podem ser explicados pelos efeitos negativos da presença de um mergulhador no comportamento de algumas espécies de peixes, em combinação com a visibilidade da água, um outro fator limitante, que influencia mais a deteção do SDOV que do SBRUV. De outro lado, a presença de isco no SBRUV permitiu atrair mais espécies, incuindo os de tamanhos grandes, para perto das câmaras, resultando numa maior riqueza e biomassa. Além destes resultados, o SDOV apresentou custos mais altos que SBRUV, devido à possibilidade de obter um número maior de amostras de SBRUV por dia. Uma comparação com UVC ainda tem que ser considerada para selecionar o melhor método não extrativo para monitorização. O UVC tem um potencial de amostrar uma comunidade de peixes mais ampla que os métodos de vídeo, devido à maior capacidade do olho humano para detetar e identificar espécies crípticas, mas é mais dependente do treino do observador em termos de identificação e capacidade de medição, não permitindo uma validação.

O presente estudo fornece conhecimentos importantes sobre dois métodos vídeo-estéreo de amostragem subaquática da biodiversidade marinha, ainda pouco usados em águas temperadas de baixa visibilidade. Este estudo é de alta impôrtancia porque permite avaliar a qualidade e/ou eficiência da reserva de Ilhotes do Martinhal, bem como fornece uma base para sua boa gestão através de monitorização biológica. Neste contexto, os nossos resultados podem servir para reavaliar as medidas e os limites atuais de proteção em comparação com os objetivos específicos desta reserva.

Palavras chave: áreas marinhas protegidas, efeito reserva, câmaras iscadas, vídeo estéreo por mergulho, peixes demersais

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## List of Abbreviations

| BRUV | Baited Remote Underwater Video |
| :--- | :--- |
| FO | Frequency of Occurrence |
| FOV | Field of View |
| MLS | Minimum Landing Size of commercial fish species |
| MPA | Marine Protected Area |
| PNSACV | Parque Natural do Sudoeste Alentejano e Costa Vicentina |
| SCUBA | Self Contained Underwater Breathing Apparatus |
| SBRUV | Stereo Baited Remote Underwater Video |
| SDOV | Stereo Diver Operated Video |
| UVC | Underwater Visual Census |

## 1. Introduction

Marine Protected Areas (MPAs) have been increasingly designated worldwide as a tool to reverse the negative anthropogenic impacts on the ocean realm (Lubchenco et al., 2003). Habitat dredging, overfishing, pollution and ocean mining can be cited as major threats for marine and coastal habitats, their biodiversity and productivity (Dayton et al., 1995; Thrush et al., 1998; National Research Council, 2001; McCauley et al., 2015). MPAs are areas set aside, mostly for the conservation of nature, aiming to protect and/or restore marine habitats, their ecosystem services and biodiversity (National Research Council, 2001; Leenhardt et al., 2015). Commonly, MPAs also promote more sustainable fisheries and rebuilt of fished populations (Gell \& Roberts, 2003; Batista et al., 2015). Based on the strictness of protection measures, a large variety of MPAs exists. Those include both 'light' or 'strong' partially protected areas, where some level of commercial and/or recreational extraction activities is allowed, and 'marine reserves', which are fully protected areas, also called 'no-take' zones, with all extraction activities prohibited (Lester \& Halpern, 2008; Lubchenco \& Grorud-Colvert, 2015). Multiple-use MPAs are designed as a combination of zones with different levels of protection, so they can meet the needs of distinct stakeholder groups (Kelleher \& Kenchington, 1992; Lester \& Halpern, 2008). Nevertheless, the implementation of protection measures and zoning often faces opposition from locals, as it contradicts the tradition of open access to the sea and generates feelings of inequalities between various groups of stakeholders when the new rights are attributed (National Research Council, 2001).
Fisheries benefits are expected from well designed and managed MPAs addressing that goal, not only in partially protected areas, but also in areas surrounding marine reserves (Gell \& Roberts, 2003; Halpern \& Warner, 2003; Giakoumi et al. 2017). Thus, fishing grounds adjacent to marine reserves might get replenished through export of fish, at different life stages, from the reserves (Russ et al., 2004; Christie et al., 2010; Harrison et al., 2012; da Silva et al., 2015). The success of MPAs was found to be positively correlated with the level of enforcement (Jennings et al., 1996; Giakoumi et al., 2017). On the other side, lack of clearly defined conservation objectives, regulation matching them, active management and enforcement creates the risk of designating a 'paper park' only (Batista et al., 2015).
After MPA implementation, it is important to assess and monitor whether MPA objectives have been met due to the protection measures in place. MPA effects can be analysed based on biological data obtained before and after the MPA implementation (Halpern, 2003; Williams et al., 2009; Horta e Costa et al., 2013a). However, when before data are not available, the
comparison between no-fishing zones (impact) and the surrounding fished areas (control) may become the only available option (Westera et al., 2003; Miller \& Russ, 2014; Malcolm et al., 2018). Given the spatial heterogeneity of distribution of marine organisms, the effects of habitat on fish communities should also be considered, as those might get confounded with reserve effect (Roberts \& Polunin, 1991; García-Charton, 2004; Miller \& Russ, 2014). To disentangle reserve effect from both temporal and spatial variability, the 'beyond-Before After Control Impact (BACI)' approach represents the most robust method, bringing together the two above mentioned approaches while using multiple periods of time and multiple control sites (Underwood, 1992; Lincoln-Smith et al., 2006). Apart from that, studies of fish species movement patterns provide additional information on reserve efficiency in terms of reserve size adequacy (Abecasis et al., 2015; Belo et al., 2016; Di Franco et al., 2018).

MPA zoning and related fishing restrictions, size of the MPA, time since protection, fishing pressure intensity before and after protection, commercial character and mobility of protected species are all factors influencing MPA efficiency (Claudet et al., 2008; Molloy et al., 2009; Curley et al., 2013; Hilborn et al., 2014; McDermott et al., 2018). Overall, commercial species show greater responses to protection than non-target species (Harmelin et al., 1995; Micheli et al., 2004; Tetreault \& Ambrose, 2007; Claudet et al., 2008; Lester et al., 2009; Horta e Costa, 2013a; Malcolm et al., 2018). Most of the positive effects of protection, such as increase in species community richness, fish densities, sizes and biomass, have been widely reported for marine reserves (García-Rubies \& Zabala, 1990; Francour, 1994; Harmelin et al., 1995; Jennings et al., 1996; Micheli et al., 2004; Williams et al., 2009; Horta e Costa et al., 2013a; Pereira et al., 2017). Large reserves were found to provide greater commercial fish densities while small reserves have been beneficial for species with rather sedentary life style as well as for small-bodied by-catch species (Rakitin \& Kramer, 1996; Tupper \& Rudd, 2002; Claudet et al., 2008; Curley et al., 2013; McLaren et al., 2015). Older reserves tend to increase fish species diversity and enhance density and biomass effects (Russ \& Alcala, 1996; McClanahan, 2000; Micheli et al., 2004; Claudet et al., 2008; Molloy et al., 2009). The effects of partial protection remain difficult to be generalized, due to the variation in protection measures between partially protected areas (Bell, 1983; Denny \& Babcock, 2004; Lester \& Halpern, 2008; Curley et al., 2013; Giakoumi et al., 2017; Malcolm et al., 2018). Anyway, even small partially-protected areas were seen to provide efficient protection of mobile commercial species (Curley et al., 2013).

Apart from positive or no effects, negative responses to protection were also found from some studies, affecting up to one third of fish species, mainly characterized as non-commercial and/or
sedentary species (Micheli et al., 2004). Those indirect effects of protection are attributed to predation-prey relationships which cause changes in the food web (Micheli et al., 2004; Takashina et al., 2012). For example, various studies confirmed that greater predation due to protection causes decrease in sea urchin abundances (Shears \& Babcock, 2003; Micheli et al., 2005; Guidetti, 2006; Clemente et al., 2011; Fernández et al., 2016). However, few studies have demonstrated indirect protection effects in prey fish, as those are more difficult to unveil (Graham et al., 2003; Willis \& Anderson, 2003; Micheli et al., 2004). As another indirect effect of protection, changes in biological habitat complexity might also contribute to shifts in fish community structure, resulting in increase in some and decrease in other species (Micheli et al., 2005; Gil Fernandéz et al., 2016). Other negative responses to protection are related to enhanced competition that favours dominant fish predators on detriment of subordinate predator species (McClanahan, 2000). Such responses are not necessarily indicative of reserve failure but rather reflect the variability of effects in different organisms (Willis \& Anderson, 2003). Apart from that, protection might bring undesired side effects such as deterioration of fish stocks and habitats in the areas that remained opened to fishing, due to redirection and intensification of fishing effort (Himes, 2003; Hilborn et al., 2004; Stevenson et al., 2013; Vaughan, 2017; Nillos Kleiven et al., 2019). Research in this topic has focused on the concentration of fishing at the reserve border which, when intense, can cause a decline in density of some, especially less mobile species, at the reserve boundary (Rakitin \& Kramer, 1996; Murawski et al., 2005; Goñi et al., 2006; Kellner et al., 2007; Nillos Kleiven et al., 2019). Although fishing benefits from fish spill-over outside from the reserve are surely desired, estimation of spatial distribution and intensity of fishing effort and its regulation need to be part of MPA design, to prevent negative impacts (Kellner et al., 2007; Nillos Kleiven et al., 2019).

Last, but not least, a success of an MPA should be evaluated considering not only biological indicators but also taking social outcomes into account (Christie et al., 2010; Horta e Costa et al., 2013b; Rossiter \& Levine, 2014). Since MPAs address more local stressors, fishers are usually the most affected group by protection measures (Hilborn et al., 2004). In case of marine reserves, displacement of fishers from current fishing areas might induce additional costs, such as increased travel time to new fishing grounds and higher requirements on fuel (Hilborn et al., 2004; Stevenson et al., 2013; Vaughan, 2017). Fishers might also perceive a decrease in catches following the reserve implementation (Cinner et al., 2014). Studies of MPAs effects on local fishing communities' welfare rather indicate that no impacts and positive impacts are more common than negative impacts, while the poorest of fishers were identified as the most vulnerable group (Mascia et al., 2010; Cinner et al., 2014). Fishermen attitude towards MPAs
often originates from the extent of involvement in decision making in the reserve establishment (Chuenpagdee et al., 2013; Thaman et al., 2016). For this reason, a strong local community participation, especially of fishermen, since the initial phase of MPA creation process, plays a key role in future MPA success (Himes, 2003; Mackelworth et al., 2008; Gall \& Rodwell, 2016; Thaman et al., 2016).

Monitoring of ichthyofauna assemblages of MPAs can be done using traditional sampling tools, such as fishing gear or fish landing data (Rakitin \& Kramer, 1996; Williams et al., 2009; Roberts et al., 2001; Horta e Costa et al., 2013a; Pereira et al., 2017). Anyway, non-destructive sampling techniques are more appropriate, as they minimize impacts on fish stocks and habitats (Malcolm et al., 2007; Unsworth et al., 2014). So far, Underwater Visual Census (UVC) performed by trained divers with Self Contained Underwater Breathing Apparatus (SCUBA) has been the most widely applied method for assessment of fish species sizes and densities in MPAs (Jennings et al., 1996; Rakitin \& Kramer, 1996; Tupper \& Rudd, 2002; Christie, 2004; Williams et al., 2009; Vasco-Rodrigues et al., 2011; Vergés et al., 2012; Henriques et al., 2013; Horta e Costa et al., 2013a). Nevertheless, UVC is known to have various biases, related to observers' level of experience, selected methodology and changes in fish behaviour in response to the presence of divers (St. John et al., 1990; Cole, 1994; Francour et al., 1999; Williams et al., 2006; Watson \& Harvey, 2007; Pais \& Cabral, 2018). In recent years, non-invasive methods based on underwater video techniques have expanded rapidly, thanks to improved image quality and prolonged recording time, achievable at relatively low costs (Cappo et al., 2003; Mallet \& Pelletier, 2014; Unsworth et al., 2014; Bouchet \& Meeuwig, 2015; Letessier et al., 2015). Underwater video systems not only enable precise fish species identification but, when mounted as stereo-video, allow for accurate measurement of fish morphometrics (Cappo et al., 2003; Boutros et al., 2015; Letessier et al., 2015). Length measurements from stereo-video systems provide greater accuracy and precision than length estimates made by experienced divers, which is of a great importance when assessing protection effects based on fish length and/or biomass (Harvey et al., 2002). Video-techniques, when compared to UVC, enable reduction of requirements on divers' expertise and provide a permanent sample record, which can be reanalysed at any time by different observers (Cappo et al., 2003; Goetze et al., 2015).

Stereo Baited Remote Underwater Video (SBRUV) is one of the stereo-video sampling techniques mostly used in MPA effects surveys (Westera et al., 2003; Denny \& Babcock, 2004; Miller \& Russ, 2014; McLaren et al., 2015; Malcolm et al., 2018). Due to the presence of a bait plume, SBRUV becomes a point of attraction for fish, thus being able to detect greater richness
of fish species and their densities than do un-baited remote video systems (Cappo et al., 2003; Hardinge et al., 2013). Higher detection and abundances of predatory and scavenger fish species, which are the groups targeted by fisheries, were reported for SBRUVs when compared to un-baited cameras, UVC and Stereo Diver Operated Video (SDOV) (Harvey et al., 2007; Goetze et al., 2015). SBRUVs favour sampling of larger bodied individuals that, through agonistic behaviour, tend to exclude smaller sized individuals from the screen view (Hardinge et al., 2013; Dunlop et al., 2015). As a result, an underestimation of small individuals might lead to bias in species length-frequencies and densities, especially in areas where fish densities are high (Dunlop et al., 2015). Apart from that, fish densities derived from SBRUV sampling are further limited, as the area of bait attraction for fish is hard to model, being influenced by variables such as currents' direction and velocity (Taylor et al., 2013; Haggitt et al., 2014). However, when compared to UVC, SBRUV is more suitable for repetitive sampling at depths above 20 meters where the complexity of dives increases (Terres et al., 2015). It also eliminates both positive or negative behavioural responses in fish to divers (Willis \& Babcock, 2000; Assis et al., 2013). However, higher richness and abundances of fish species have been associated to UVC rather than SBRUV, due to divers' greater ability to detect both cryptic, crypto-benthic, herbivorous and territorial fish species (Langlois et al., 2006; Stobart et al., 2007; Colton \& Swearer, 2010; Lowry et al., 2012), although greater abundances were reported from SBRUV than UVC for specific groups of species (Willis et al., 2000; Willis \& Babcock, 2000; Goetze et al., 2015). Thus, each technique might be more suitable for sampling different fish species and result in different assemblage structure (Willis \& Babcock, 2000; Colton \& Swearer, 2010; Goetze et al., 2015).

SDOV represents another stereo-video technique that has been used in assessment of fish species composition, densities and sizes, although its application as an MPA monitoring tool remains scarce (Watson et al., 2005; Tessier et al., 2013; Goetze et al., 2015; Wartenberg \& Booth, 2015; Navarro-Martínez et al., 2017). SDOV tends to provide lower fish species richness and abundances than UVC, explained by greater detection capacity of the human eye when compared to the cameras (Pelletier et al., 2011; Holmes et al., 2013; Tessier et al., 2013). When combined with the behavioural response in fish to divers and effects of bait, lower species richness and abundances were reported from SDOV than from SBRUV (Watson et al., 2005; Langlois et al., 2010; Watson et al., 2010). However, SDOV has a potential as a monitoring tool for easily identifiable, non-shy and/or herbivorous target fish species (Tessier et al., 2013; Goetze et al. 2015). As a complementary method to SBRUV, SDOV avoids alterations in fish behaviour due to the bait plume while it also allows permanent recording.

Both mentioned stereo-video methods share several limitations. First, not all fish always appears in the field of view (FOV) of both cameras, which excludes them from length measurement, resulting in narrower length datasets and lower statistical power when compared to UVC (Holmes et al., 2013; Cundy et al., 2017; Wilson et al., 2018). In addition, most of the studies that used stereo-video so far, took place in tropical waters that are characterized by good visibility conditions. In temperate waters, light conditions and visibility are limiting factors, causing lower video quality and shortening the distance in FOV, again causing a decrease in the proportion of fish that can be precisely measured (Unsworth, 2014).

Furthermore, processing of video imagery is known to be time consuming. Apart from costs of the camera equipment, laboratory time represents net extra costs of SBRUV and SDOV above UVC (Holmes et al., 2013; Tessier et al., 2013). However, depending on the amount of data gathered from stereo-video, time expenditures can become comparable to those of UVC (Goetze et al. 2015). SBRUV was found to be more cost-effective than SDOV, as staff time per sampling site reduces when several SBRUV frames are deployed simultaneously (Langlois et al., 2010). Thus, SBRUV might become a more competitive option to UVC. In fact, combination of sampling methods is desirable as it will help to understand limitations of each method and gather complementary results for the sampled area, increasing analysis robustness (Cappo et al., 2003; Watson et al., 2005).

In this study, we aimed at assessing whether potential effects of protection existed in the marine reserve of Ilhotes do Martinhal, which is part of a large MPA of the South-West coast of Portugal. The protection effect in demersal fish and commercial invertebrate community was evaluated based on richness, abundance, length and biomass. Response to protection was expected to be found in target (commercial) species and especially in target species above minimum landing size (MLS), which are the groups most sensitive to prohibition of fishing. Given the lack of data on demersal fish and commercial invertebrates densities and sizes from the time prior to reserve implementation, a control-impact approach between reserve inside and outside was applied. As protection effects might be confounded with effects of habitat, we also checked for influence of physical habitat. SBRUV was used as a primary sampling method, as previous studies (Capaz, 2013; Willenbrink, 2016) confirmed that the method performed well in Portuguese coastal waters. SDOV was added as a complementary sampling method to enable for SBRUV results validation and complementarity. At the same time, the sampling performance of SDOV was tested as this method is rather pioneer in temperate waters. As both selected methods substantially reduce requirements on observer's in situ expertise, both have a potential as easily applicable reserve monitoring tool. Sampling performance and cost-
effectiveness were compared between methods to understand if those would be suitable for future reserve monitoring.

## 2 Materials and methods

### 2.1 Study area

The "Parque Natural do Sudoeste Alentejano e Costa Vicentina (PNSACV)" is a marine protected area (MPA) that was established in 1995 along the south-west coast of Portugal (Decreto Lei $n^{\circ}{ }^{\circ} 26 / 95$ ). With 2 km of width, the MPA spreads continuously for 130 km of the Portuguese coast, between Sines and Burgau (Figure 1). Various types of marine ecosystems can be found inside the MPA, given the presence of rocky and sandy bottoms and small estuaries. In 2011, with the approval of the management plan of the marine park, the MPA was divided into zones with different levels of protection: full protection (PT), partial protection of type I (PPI) and type II (PPII) and a complementary protection or buffer zone (Resolução do Conselho de Ministros $n .^{\circ}$ 11-B/2011). All types of fishing and other human activities have been prohibited inside of the PT zone. Nine rocky islets, with surrounding distance up to 100 m , were assigned this category. Except for non-extractive recreational activities, allowed within partial protection zones, fishing is forbidden in PPI, except for commercial hand harvest of goose barnacles (Pollicipes pollicipes) in the coastal cliffs. Thus, the PPI represents a 'marine reserve' (also known as no-take) for fish and invertebrates (except for the intertidal species $P$. pollicipes). Four locations with rocky habitat and sand belong to the PPI category, including Ilhotes do Martinhal, which is the reserve of this study (Figure 1).

The buffer zone covers the majority of PNSACV MPA. In this zone, commercial fishing can only be conducted by locally registered boats with a valid license for PNSACV. There are several restrictions of fishing gear usage, but they are derived from the national law and thus are not MPA specific. In this context, purse seines are allowed further than 0.25 nm from coast with a minimum depth of 20 meters, same as defined in currently applicable national legislation (Portaria n. ${ }^{\circ}$ 1102-G/2000; Resolução do Conselho de Ministros n. ${ }^{\circ}$ 11-B/2011). The only additional limitation to fishing gear types in relation to the national legislation is for longlines that can only be deployed further than 0.5 nautical miles ( nm ) from coast instead of nearshore (Portaria n. ${ }^{\circ}$ 1102-C/2000; Resolução do Conselho de Ministros n. ${ }^{\circ}$ 11-B/2011). Recreational fishing (hand harvest, handlining and spearfishing) have been allowed in the buffer zone within the scope of national legislation, except when specific legislation exists, such as temporal closures for particular fish species and lower daily limits for both fishing and hand harvest
(Portaria n. ${ }^{\circ}$ 115-A/2011; Decreto-Lei $n .{ }^{\circ}$ 101/2013). For land-based commercial and recreational handline fishers, the capture of Diplodus sargus and Diplodus vulgaris has been forbidden between February $1^{\text {st }}$ and March $15^{\text {st }}$. Between March $1^{\text {st }}$ and May $31^{\text {st }}$ the same restriction has been valid for Labrus bergylta (Portaria n. ${ }^{\circ}$ 115-A/2011; Portaria n. ${ }^{\circ}$ 115$B / 2011$ ). In the whole MPA, it has been strictly prohibited to capture Epinephelus marginatus, marine mammals, marine and/or migratory birds. Both commercial fishing with sets of octopus traps and land-based handlining represent fishing activities that can be commonly observed around Atalaia and Barranco, which are part of the buffer zone assessed in the current study.

This study was held in the south coast of PNSACV in Ilhotes do Martinhal 'marine reserve' which covers an area of $3.9 \mathrm{~km}^{2}$ with rocky and sandy bottom. Multiple locations with rocky habitat from both inside (i.e. from the PPI zone) and outside the marine reserve (i.e. from the buffer zone) were sampled. Sampling locations outside the marine reserve were located both in the west and in the east of the reserve (Figure 1).


Figure 1: Map of PNSACV MPA and Ilhotes do Martinhal marine reserve (Partial Protection I). A) Map of the PNSACV MPA in the South-West coast of Portugal, showing nine fully protected locations, four locations with Partial Protection I and the buffer zone. Full protection: I) Pedra do Burrinho, II) Pedras da Ilha do Pessegueiro, III) Pedra da Atalaia, IV) Pedra da Enseada do Santoleiro, V) Pedra de Baía da Nau, VI) Pedra da Carraca, VII) Pedra da Agulha, VIII) Pedra das Gaivotas, IX) Pedra do Gigante. Partial protection (PPI):1) Ilha do Pessegueiro, 2) Cabo Sardão, 3) Rogil, 4) Ilhotes do Martinhal. B) Map of sampling locations inside of Ilhotes do Martinhal marine reserve and outside, including three sampling locations inside (Martinhal, Ponta dos Caminhos, Barranco inside) and four sampling locations outside (west outside: Grutas, Atalaia; east outside: Barranco outside, Ingrina).

### 2.2 Sampling methods

### 2.2.1 SBRUV sampling design

The SBRUV systems used in this study were developed by Willenbrink (2016) as a modification of a single camera BRUV build by Capaz (2013). Each SBRUV consisted of two small action cameras in waterproof housings, mounted at 0.4 m distance and facing the bait basket at $8^{\circ}$ inward angle (Figure 2 and 3). Three SBRUVs were used for each sampling site, one with two GoPro Hero3 cameras and two with small action cameras (SK8 CAM 4K), providing low cost sampling option at acceptable image quality (Letessier et al., 2015). All cameras were set with a medium FOV of $94.4^{\circ}$ in air, corresponding to $67^{\circ} \mathrm{FOV}$ in seawater. All videos were recorded in full HD quality ( $1080 \times 25$ frames for GoPro, $1080 \times 30$ frames for SK8). Considering 3 m as average visibility of the video recordings, the common FOV of the SBRUV systems was estimated to be $5.15 \mathrm{~m}^{2}$ with 3 m of distance ahead of the cameras, counted in GeoGebra software (International GeoGebra Institute, 2018) (Figure 3).


Figure 1: Sketch of SBRUV and SDOV systems. Image created with Paint 3D.


Figure 3: Field of view (FOV) of SBRUV and SDOV. Upper figure: SBRUV, lower figure: SDOV, L: left camera, R: right camera, LR: distance between left and right camera, IJ : width of the common FOV at 3 m of distance, MN: width of total FOV at 3 m of distance, SHJIG: area of common FOV, SRHNMGL: area of total FOV, $\alpha$ : inward angle of each camera, $\beta$ : FOV angle of each camera in seawater, $\delta$ : common FOV angle in seawater. Image made in GeoGebra software.

Three SBRUV sampling campaigns took place between March and May 2018. During each campaign, seven locations, three inside the reserve and four outside, were sampled (Figure 1). At each location, the three SBRUV systems were deployed from a boat at rocky bottom habitats of the infralittoral zone (from 7 to 18 m of depth). A minimum distance of 250 meters was kept between each two SBRUVs to ensure independence of samples, by minimising the likelihood that fish can swim between two SBRUVs during sampling (Cappo et al., 2001; Langlois et al., 2010; Haggitt et al., 2014; Goetze et al., 2015). This minimum distance was defined considering the total available area of rocky habitats relative to replication requirements while assuming
that bait plume attraction power was lower than 250 m and that demersal fish species targeted by our sampling were reef dependent, thus showing reduced mobility. The net time of one SBRUV recording was 30 minutes, as it was confirmed to be the enough time to reach maximum demersal fish species richness (Capaz, 2013). Sampling was performed during daylight hours, between 9 am and 17 pm , to avoid variability between samples due to fish crepuscule behaviour (Harvey et al., 2007). The same total quantity ( $\sim 200 \mathrm{~g}$ ) and proportions of small pelagic fish (Sardina pilchardus, Trachurus trachurus, Scomber colias) and mussels (Mytilus galloprovincialis) were used as bait throughout the experiment (Willis \& Babcock, 2000; Capaz, 2013). Bait basket content was changed after every two SBRUV deployments.

On a second day of each sampling campaign, resampling for excluded samples (see reasons below) took place to maintain the requirements on the number of replicates as close as possible to the initially planned design. As initially planned, 63 samples were collected with SBRUVs, but out of those, 15 samples were excluded ( $23.8 \%$ ). The reasons for samples exclusion were as follows: system landed on sandy bottom ( $53.3 \%$ of excluded samples), water visibility conditions were less than $2 \mathrm{~m}(20.0 \%)$, system landed with obstructed FOV ( $20.0 \%$ ) and camera's FOV changed compared to the calibrated image ( $6.7 \%$ ). Resampling was then held for 12 (out of the 15) excluded samples and had a $100 \%$ success rate, as the main reason of failure was landing on sand, which was resolved by sampling point correction. The other three excluded samples were rejected ex-post during data analysis (due to bad water visibility and camera's FOV change), and thus not subject to resampling. A final number of 60 valid samples (26 inside and 34 outside the marine reserve, in line with the planned 3:4 inside vs. outside ratio) was used in data analysis.

### 2.2.2 SDOV sampling design

The SDOV system was derived from the SBRUV (Figure 2) but cameras had an inward angle of $15^{\circ}$ (Figure 3). Three SDOV sampling campaigns took place between April and July 2018. Two locations inside the marine reserve and two to three locations outside (both in the west and in the east) were sampled during each campaign (Figure 1). The SDOV fish census was performed using SCUBA diving, with one diver stretching a transect line and a second diver swimming at his side and video recording. At each location, 30 m transects were sampled (Schmitter-Soto et al., 2018). Distance of 10 m was left between the transects (Goetze et al., 2015). At each location, three replicate transects were sampled during each campaign (Monteiro et al., 2012). Divers swam no more than one meter above the substrate at a constant speed of
approximately $8 \mathrm{~m} / \mathrm{min}^{-1}$ (Lincoln Smith, 1988; Cheal \& Thompson, 1997). The total width of the transect was calculated to be 6.4 m at 3 m ahead the SDOV system (Figure 3), providing a total transect area of $192 \mathrm{~m}^{2}$ (area where demersal taxa was counted). When recording, cameras faced forward and slightly down, with the aim to capture demersal fish species (Holmes et al., 2013; Wilson et al., 2018). However, when fish seemed to have appeared in the FOV of only one camera, the cameras were turned towards the fish as this was found to be useful for video post-processing (length measurement). The SK8 cameras were used for SDOV sampling as they allowed the recorded video to be visualized directly in the cameras screen when recording, thus helping to control the position of the cameras relative to the bottom. A total of 39 samples (18 inside and 21 outside) were obtained with the SDOV and used in the analysis.

### 2.2.3 Calibration procedure

Calibration of both stereo-video systems followed the procedure of Neuswanger et al. (2016), including video synchronisation, distortion correction and 3D calibration. On top of that, cameras and housings were labelled, allowing the same housing to be used with the same camera throughout the whole experiment. Such settings enabled the use of one distortion correction frame for each camera during the entire study, thus reducing the time and costs of video post-processing. At the beginning of each sampling day, housings with cameras were tightly screwed into their base on the stereo-video structure, to avoid further housing movements during sampling. Hands were clapped at the beginning of each recording for synchronisation purposes of each pair of cameras.

### 2.3 Data processing and analysis

### 2.3.1 Stereo-video analysis

Processing of video material was done in VidSync software version 1.661 (Neuswanger et al., 2016). The processing included i) merge and synchronisation of the two videos, distortion correction and 3D calibration, ii) species abundance counts, iii) individual length measurements and iv) habitat categorization. For the SBRUV, MaxN, which is the maximum number of individuals of a species present in a frame or during an interval of time, within the 30 min video recording, was used as count of relative abundance (Cappo et al., 2003). This technique is currently considered as the best proxy for fish relative abundance obtained from SBRUVs, while it avoids repeated counts of the same individuals. On the other side, MaxN is a conservative measure, which might lead to underestimation of the actual fish abundance (Cappo
et al., 2003; Watson et al., 2005). Taking advantage of the video technique, fish were counted for 30 seconds which were added to the MaxN frame, while making sure that individuals were not counted twice (Stobart et al., 2007; Willenbrink, 2016). MaxN counts included individuals that appeared only in one of the two videos (total FOV), whereas length measurements could only be done when individuals appeared in both videos (common FOV; Figure 3). For each species, within its respective MaxN frame, total length (cm) was measured in fish and mantle length in cephalopods (Horta e Costa et al., 2013a; Unsworth et al., 2014; Wilson et al., 2018). Habitat category was assigned to each SBRUV sample based on visual assessment of physical habitat complexity from the videos. Two criteria, surface type and abundance of hidings, were considered when deciding on habitat category (Table 1). In addition to the identification of species, their respective abundance and size, and overall habitat features, the presence of illegal fishing gear inside the marine reserve was noted down when encountered during field-sampling or in video footages.

Table 1: Physical habitat category of SBRUV samples.

| Criteria/ Habitat | hab1 | hab2 | hab3 | hab4 |
| :---: | :---: | :---: | :---: | :---: |
| Criteria 1: <br> surface type | flat | flat to irregular | irregular to highly <br> irregular | highly irregular |
| Criteria 2: <br> abundance of hidings | very few | few | some | common |
| Description | rock mixed <br> with sand, flat <br> mother rock | moderately <br> structured mother <br> rock | highly structured <br> mother rock or <br> mixed with small <br> boulders | big boulders |

SDOV video analysis followed the same steps as SBRUV analysis, except that instead of using MaxN as abundance estimate, demersal fish and commercial invertebrates were counted along the whole transect, except for fish that appeared from behind of the diver (Goetze et al., 2015). Habitat category was not tested for SDOV samples.

### 2.3.2 Data processing

Both selected stereo-video methods were designed to sample demersal species of fish and invertebrates (cephalopods). Schools of semi-pelagic fish (Boops boops) were excluded from the analysis due to their underestimation by this technique targeting species highly associated to the rocky bottom, and to their little potential to provide a response to protection (Horta e Costa et al., 2013a). In addition, cryptobenthic species, such as Gobiidae or Blenniidae, were
considered out of scope as the horizontal video also underestimates those taxa (Watson et al., 2005; Holmes et al., 2013).

The number of sampled demersal fish and invertebrate species provided information on species richness that was expressed by sample (i.e. richness per sample) and by protection (i.e. total richness). Frequency of occurrence (FO) was expressed for each demersal species, based on the equation:
$\mathrm{FO}(\%)=\left(\mathrm{N}^{\circ}\right.$ of samples containing the species/ Total $\mathrm{N}^{\circ}$ of samples $) * 100$

Biomass was calculated for every individual based on length measurements in MaxN frames, using species-specific length-weight relationships (Gonçalves et al., 1997; Borges et al., 2000; Morato et al., 2001; Morey et al., 2003; Sifner \& Vrgoc, 2004; Veiga et al., 2009; Horta e Costa et al., 2013a). When not available, FishBase species information from closest regions (and with enough individuals tested) was used (Froese \& Pauly, 2019). For each sample, mean species biomass was multiplied by species abundance (MaxN for SBRUV, total count for SDOV) to obtain species biomass per sample (Willis et al., 2003; Willenbrink, 2016). Total biomass was then calculated as the sum of biomass of all species in a sample. In the same way, total abundance was obtained as the sum of abundance of all species in a sample.

Identified taxa were grouped based on their local fishing commercial status as target (commercial) or non-target (non-commercial). Target species were then grouped according to their size in relation to minimum landing size (MLS), when available. Commercial species without MLS defined were not classified as neither above nor below MLS.

### 2.3.3 Statistical analysis

SBRUV: The effects of factors protection (two levels: 'inside' vs. 'outside' marine reserve) and habitat (four levels: hab1, hab2, hab3, hab4) on response variables of richness, abundance, length and biomass were tested. The following community datasets for abundances (MaxN) and biomass (at the MaxN) were created: all demersal species, target species, target species above MLS, target species below MLS and non-target species. Primer-E version 6.4.7.0 with PERMANOVA+ package was used for multivariate analysis (Anderson, 2005). First, a square root transformation was applied to species abundance and biomass by sample, to increase the contribution of less dominant species (Clarke \& Madwick, 2015). Two-way crossed main PERMANOVA with 9999 permutations was conducted based on Bray-Curtis similarity matrix. For habitat, pair-wise PERMANOVA was conducted only when main PERMANOVA test was
significant for that factor ( $\mathrm{p}<0.05$ ). In addition, SIMPER analysis was done to understand which species contributed the most to differences between protection levels (Clarke \& Madwick, 2015).
Univariate analysis was done using R version 3.4.3 (R Core Team, 2017). For community datasets, effects of protection and habitat on species richness, total abundance and total biomass per sample were tested. At individual species level, differences in abundance, length and biomass per sample were inspected for the key-target demersal species, Diplodus sargus and Diplodus vulgaris, and for the most common non-target species, Coris julis, Serranus cabrilla and Labrus bergylta. For the key-target species, we also analysed whether differences in abundance and biomass existed for individuals above MLS and below MLS. All datasets were first tested for normal distribution (Shapiro-Wilk test) and homogeneity of variances (Bartlett's test) (Dytham, 2011; Gardener, 2017). As none of the datasets complied with conditions of parametric tests, differences in response variables based on protection and habitat were tested using non-parametric statistical tests (Wilcoxon rank-sum test, Kruskal-Wallis test) (Montgomery \& Runger, 2003; Dytham, 2011; Gardener, 2017). For explanatory purposes, we also tested whether differences in response variables existed between the west outside, the east outside and the reserve inside. Univariate analysis was accompanied by descriptive statistics (mean $\pm$ standard error).

To investigate whether habitat was independent from protection, we used Chi-square Test of Independence with simulated p-value (Dytham, 2011; Montgomery \& Runger, 2003). Furthermore, interaction between protection and habitat for abundance and biomass was tested in multivariate analysis (two way crossed PERMANOVA) (Anderson, 2005).

SDOV: Statistical analysis of the SDOV data pursued the same scheme as for SBRUV with the following differences: i) factor habitat was not available and thus not tested, ii) abundances and biomass data were based on total counts and length measurements obtained along each transect and not on MaxN.

### 2.3.4 Methods comparison

Costs of the two-stereo video methods were compared using variable and fixed costs. Variable costs per sample were calculated from fieldwork costs of day of sampling and from laboratory costs. Fieldwork costs included costs of travel to Sagres and back, boat expenses, labour costs and price of consumables specific to each method during one sampling day (Perkins et al., 2013). Laboratory costs included labour costs of data post-processing (data storage, merging of
automatically split videos, calibration in VidSync, abundance counts and length measurements). Both fieldwork and laboratory costs were recounted to costs per one sample. Labour costs were based on average daily income of a research technician in Portugal in 2018 ( $48 € /$ day). Fixed costs were expressed as 'initial set up costs’, including the price of all equipment needed to be able to start using the method (Perkins et al., 2013). Those involved diving material for SDOV and stereo-video material for both SDOV and SBRUV.

Using univariate statistical methods (Wilcoxon rank-sum test), we assessed whether differences existed between methods in sampled values of richness, abundance, length and biomass (Montgomery \& Runger, 2003; Dytham, 2011; Gardener, 2017). We also compared sampling performance between methods based on the number of valid samples collected during one day of sampling. Taking into consideration that not all individuals can always be measured from stereo-video recordings, the percentage of measured out of all counted (sum of MaxN in SBRUV, total count in SDOV) individuals of a species was calculated for the most common species, to compare methods efficiency in obtaining fish sizes.

## 3 Results

### 3.1 SBRUV

### 3.1.1 Frequency of occurrence, species richness and illegal fishing gear

A total of 30 taxa from 15 different families were identified from SBRUV samples (Appendix, Table 12). Most of the taxa belonged to Sparidae ( 9 species) and Labridae ( 6 species) families. Out of all, 19 taxa were classified as target and 11 as non-target species. Individuals and schools of $B$. boops were repeatedly spotted but excluded from the analysis due to their semi-pelagic character. Apart from demersal fish species, three target species of Cephalopoda (O. vulgaris, L. vulgaris, S. officinalis) were also present. From target species, the highest frequency of occurrence was related to D. sargus ( 96.7 \% of the samples) and D. vulgaris ( $95.0 \%$ ). From non-target species, C. julis ( $98.3 \%$ ) and S. cabrilla (81.7\%) were the most common. A total of 22 taxa appeared inside the marine reserve and 26 outside ( 21 in west outside and 23 in east outside). There were four species only spotted inside, three of high commercial value (Pagrus pagrus, Pagrus auriga, L. vulgaris). Other 8 species only appeared outside the reserve, corresponding to both target (S. officinalis, Conger conger, Mullus surmuletus, Trisopterus luscus, Sparus aurata) and non-target species (Chromis chromis, Pseudocaranx dentex, Serranus atricauda) (Appendix, Table 12).

Mean richness was found to be significantly higher inside than outside (Wilcoxon test, $\mathrm{p}=0.012$ ). Mean richness by protection level was equal to $7.42 \pm 0.40$ taxa inside the reserve and $6.13 \pm 0.30$ taxa outside. This difference was due to lower mean richness in east outside when compared to the reserve inside (Wilcoxon test, $\mathrm{p}=0.015$, Appendix, Table 13). Those results were obtained despite the total richness in east outside (23 taxa) and inside (22 taxa) were similar. When looking at mean richness in habitats, no differences were detected (Kruskal test, $p=0.176$, Figure 4).

Illegal fishing gear, represented by sets of octopus traps, was encountered on two occasions during SBRUV sampling campaigns. Both encounters occurred at Barranco inside location (on 27/03/2018 and on 24/05/2018). As each campaign covered two sampling days, this means that illegal fishing gear was found two out of six days ( $\sim 33 \%$ of sampling days). During a later sampling campaign, octopus traps were found again at Martinhal location (18/07/2018, biological data of this sampling campaign not included in this study).


Figure 4: Boxplot of richness by protection and by habitat (SBRUV). Significant differences marked with * when inside>outside.

### 3.1.2 Results for community datasets

## Effect of protection

Multivariate analysis showed significant differences in abundance of all demersal species between inside and outside the reserve (PERMANOVA; Table 2), with higher average values observed for the reserve in most species (SIMPER; Appendix, Table 16). For all other community datasets (target species, target species above MLS, target species below MLS and non-target species), differences between protection levels in multivariate abundance were not confirmed, although result for target species was marginally non-significant (PERMANOVA,
$\mathrm{p}=0.06$; Table 2). Differences in abundance between protection levels (average dissimilarity of all demersal species group: 43.3 \%) were mainly attributed to C. julis ( $17.5 \%$ contribution to dissimilarities), D. vulgaris (10.1 \%), D. sargus (10.0 \%), S. cabrilla (6.4 \%), Mugilidae (5.8 $\%$ ) and L. bergylta ( $5.7 \%$ ) (SIMPER; Appendix, Table 16). Average abundances of all species, except for D. vulgaris, Mugilidae and Sarpa salpa, were slightly higher inside than outside, thus indicating that a trend of greater abundances inside was present (Appendix, Table 16). The species that showed more positive results in the reserve were, from target species: D. sargus, Oblada melanura, Diplodus cervinus and $O$. vulgaris, and from non-target species: C. julis, S. cabrilla, L. bergylta and Ctenolabrus rupestris.

For total abundance, univariate analysis did not show significant differences between protection levels (inside vs. outside) in any of the tested community datasets (Table 2). However, significant differences in total abundance existed when comparing the reserve to the west outside and to the east outside separately. For all demersal species and target species, total abundance was significantly greater in west outside than inside the reserve, and inside the reserve than in east outside. For non-target species, total abundance was also greater inside than in east outside (Appendix, Table 13).

Table 2: Differences in community abundance and biomass between protection levels (SBRUV). Results of PERMANOVA and Wilcoxon test, significant p-values ( $\mathrm{p}<0.05$ ) marked *. For Wilcoxon test: inside>outside marked * and outside <inside marked ${ }^{* *}$. MLS: minimum landing size.

|  | PROTECTION (SBRUV) |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
|  | PERMANOVA |  | Wilcoxon test |  |
| Community dataset | Abundance | Biomass | Total abundance | Total biomass |
| All demersal species | $0.003^{*}$ | $0.001^{*}$ | 0.99 | $0.01^{*}$ |
| Target species | 0.06 | $0.003^{*}$ | 0.77 | $0.001^{*}$ |
| Target species above MLS | 0.07 | $0.03^{*}$ | 0.35 | $0.002^{*}$ |
| Target species below MLS | 0.16 | 0.43 | 0.84 | 0.16 |
| Non-target species | 0.10 | $0.03^{*}$ | 0.56 | $0.01^{*}$ |

Multivariate results (PERMANOVA) showed that biomass of all community datasets, except target demersal species below MLS, differed significantly between reserve inside and outside (Table 2). Larger significant differences were found in all demersal species and in the target species group (PERMANOVA, p < 0.01). Differences in biomass between protection levels (average dissimilarity in all demersal species: $59.2 \%$ ) were associated with D. sargus ( $14.3 \%$ contribution to dissimilarities), L. bergylta ( 8.7 \%) D. cervinus ( 8.6 \%), Mugilidae ( $8.3 \%$ ) and
D. vulgaris (7.6 \%) (SIMPER; Appendix, Table 16). Except D. vulgaris, Mugilidae and S. salpa, higher average biomass inside than outside was found in all other species. Out of those, the ones that seemed to have thrived the most inside the reserve were, from target taxa, $D$. sargus, $D$. cervinus, $O$. melanura and $O$. vulgaris and, from non-target taxa, L. bergylta and $C$. julis (SIMPER; Appendix, Table 16). Although biomass of Dicentrarchus labrax and P. auriga had a small but non-negligible contribution to dissimilarities of target species ( $7.2 \%$ and $4.2 \%$ respectively), the rare character made them less relevant as indicators of differences (SIMPER; diss $/ \mathrm{SD}=0.43$ and diss/ $\mathrm{SD}=0.53$; Appendix, Table 16).

Univariate analysis confirmed the same results for total biomass as PERMANOVA, showing that for all tested community datasets, except target species below MLS, total biomass was significantly greater inside than outside the reserve (Table 2). Larger significant differences were found in the datasets of target species and target species above MLS (Wilcoxon test, $\mathrm{p}<$ $0.01)$. When comparing the reserve with west and east outside, total biomass was found significantly greater inside the reserve than in the east outside, for all tested datasets (Wilcoxon test, $\mathrm{p}<0.05$; Appendix, Table 13). Only for target species above MLS, total biomass was also significantly greater inside the reserve than in west outside (Wilcoxon test, $\mathrm{p}<0.05$ ). Total biomass of community assemblages is displayed in Figure 5.


Figure 5: Total biomass of community datasets by protection (SBRUV). Mean $\pm$ standard error displayed. ADS: all demersal species, TS: target species, TS+: target species above MLS, TS-: target species below MLS, NTS: non-target species. Significant differences marked $*$ when inside>outside and ${ }^{* *}$ when inside<outside. MLS: minimum landing size.

## Effect of habitat

Multivariate analysis indicated that abundance of all demersal species and of non-target species differed significantly between habitats (PERMANOVA, $\mathrm{p} \leq 0.5$; Table 3). Based on pair-wise PERMANOVA, significant differences in abundance of all demersal species and non-target species were found between hab1, the less complex habitat, and other habitats (Table 3). Differences in abundance between hab1 and other habitats (average dissimilarity of all demersal species between 43.9 \% and 46.4 \%) was accredited to $C$. julis (> $18.9 \%$ contribution), $D$. sargus (> $8.4 \%$ ), D. vulgaris (> 9.1 \%), S. cabrilla (> 6.6 \%) and C. rupestris (> 5.6 \%) (SIMPER, Appendix; Table 17). Average abundance was higher in hab1 than in all other habitats for $D$. sargus but lower for the other four above stated species. From non-target species, C. rupestris was absent from hab1 (SIMPER). No differences between habitats were detected for biomass based on multivariate analysis (PERMANOVA, $\mathrm{p}>0.05$; Table 3), with the result for all demersal species being marginally non-significant (PERMANOVA, $\mathrm{p}=0.068$ ). Univariate analysis did not reveal any significant differences between habitats neither for total abundance nor total biomass (Kruskal-Wallis test, $\mathrm{p}>0.05$; Table 3).

Table 3: Differences in community abundance and biomass between habitats (SBRUV). Results of PERMANOVA and Kruskal-Wallis test, significant p-values ( $\mathrm{p}<0.05$ ) marked with *. For pairwise PERMANOVA, only significant differences are shown. MLS: minimum landing size.

|  | HABITAT |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
|  | PERMANOVA |  |  | Kruskal-Wallis test |  |
| Community dataset | Abundance |  | Biomass | Total <br> abundance | Total <br> biomass |
| All demersal species | $0.03^{*}$ | Hab1 x Hab3 <br> Hab1 x Hab4 | 0.07 | 0.31 | 0.41 |
| Target species | 0.18 | - | 0.11 | 0.13 | 0.30 |
| Target species above MLS | 0.62 | - | 0.11 | 0.55 | 0.60 |
| Target species below MLS | 0.88 | - | 0.08 | 0.23 | 0.62 |
| Non-target species | $0.05^{*}$ | Hab1 x Hab2 <br> Hab1 x Hab3 <br> Hab1 x Hab4 | 0.13 | 0.10 | 0.98 |

## Interaction between protection and habitat

For the whole area and for the reserve inside, hab3 had the lowest proportion ( $18 \%$ ) of all habitats (Figure 6). In addition, habitats with lower complexity had greater proportions inside the reserve (inside: hab1=35 \%, hab2=31 \%; outside: hab1=21\%, hab2= $21 \%$ ). Outside the reserve, hab4 had the highest proportion (inside: $23 \%$; outside: $35 \%$ ) while the other habitats
had equal or similar proportions (Figure 6). There was no significant relationship found between the two factors, protection and habitat (Chi-square Test of Independence, $\mathrm{p}=0.33$ ). No significant interaction between protection and habitat existed neither for abundance, nor biomass in multivariate analysis (PERMANOVA, p>0.05) (Table 4).


Figure 6: Proportions of habitat categories in the whole sampled area and by protection.

Table 4: Results of PERMANOVA for interaction between protection and habitat (SBRUV). MLS: minimum landing size.

| Factor | PROTECTION x HABITAT |  |
| :---: | :---: | :---: |
| Statistic | Main PERMANOVA |  |
| Community assemblage/ Variable | Abundance | Biomass |
| All demersal species | 0.54 | 0.49 |
| Target species | 0.73 | 0.59 |
| Target species above MLS | 0.83 | 0.68 |
| Target species below MLS | 0.72 | 0.12 |
| Non-target species | 0.34 | 0.42 |

### 3.1.3 Results for key target species

## Effect of protection

No protection effect was found in abundance of D. sargus, D. sargus above MLS nor below MLS (Wilcoxon test, p>0.05; Table 5), although for D. sargus above MLS the result may be considered marginally non-significant (Wilcoxon test, $\mathrm{p}=0.065$ ). Biomass of $D$. sargus was found to be significantly higher inside than outside, same as for $D$. sargus above MLS (Wilcoxon test, $\mathrm{p}<0.01$; Table 5, Figure 8 ). The results of comparison between reserve inside, west outside and east outside are displayed in Appendix, Table 13, showing that for D. sargus and $D$. sargus above MLS, the biomass was significantly greater inside reserve when compared
to each of the outside areas separately. The length of $D$. sargus was confirmed to be significantly greater inside than outside (Wilcoxon test, $\mathrm{p}=0.012$; Table 5). Individuals of 1520 cm class prevailed above other length classes both inside and outside. The length-frequency distribution reflected the shift of size categories toward greater sizes inside reserve when compared to outside (Figure 7).


Figure 7: Length-frequency distribution of key target species by protection (SBRUV).

Table 5: Differences in abundance, length and biomass of key target species between protection levels (SBRUV). Results of Wilcoxon test with significant p-values ( $\mathrm{p}<0.05$ ) marked with * when inside>outside and $* *$ when inside<outside. MLS: minimum landing size.

| Factor | PROTECTION |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Statistic | Wilcoxon test |  |  |  |
| Species/ Variable | Abundance | Length | Biomass |  |
| D. sargus |  |  |  |  |
| D. sargus | 0.26 | $0.012^{*}$ | $0.05^{*}$ |  |
| D. sargus above MLS | 0.065 | $\mathrm{n} / \mathrm{a}$ | $0.003^{*}$ |  |
| D. sargus below MLS | 0.85 | $\mathrm{n} / \mathrm{a}$ | 0.560 |  |
| D. vulgaris |  |  |  |  |
| D. vulgaris | 0.136 | 0.572 | $0.037^{* *}$ |  |
| D. vulgaris above MLS | $0.029^{* *}$ | $\mathrm{n} / \mathrm{a}$ | $0.025^{* *}$ |  |
| D. vulgaris below MLS | 0.25 | $\mathrm{n} / \mathrm{a}$ | 0.550 |  |



Figure 8: Biomass of key target species by protection (SBRUV). Mean $\pm$ standard error displayed. DS: D. sargus, DS+: D. sargus above MLS, DS-: D. sargus below MLS. DV: D. vulgaris, DV+: D. vulgaris above MLS, DV-: D. vulgaris below MLS. Significant differences ( $\mathrm{p}<0.05$ ) marked with * for inside>outside and ** for inside<outside. MLS: minimum landing size.

For D. vulgaris and D. vulgaris below MLS, no significant differences in abundance existed based on protection (Wilcoxon test, $\mathrm{p}>0.05$; Table 5). However, abundance of $D$. vulgaris above MLS was significantly greater outside than inside (Wilcoxon test, $\mathrm{p}=0.029$; Table 5). Biomass was found to be significantly greater outside than inside for $D$. vulgaris and $D$. vulgaris above MLS (Wilcoxon test, $\mathrm{p}<0.05$; Table 5, Figure 8). For D. vulgaris, both abundance and biomass were significantly greater in west outside than inside, except for biomass of D. vulgaris below MLS that did not show significant differences (Appendix, Table 13). Length-frequency distribution showed that individuals of $10-15 \mathrm{~cm}$ class were the most frequent both inside and outside the reserve (Figure 7). No significant differences in length were confirmed for $D$. vulgaris based on protection (Wilcoxon test, $\mathrm{p}>0.05$, Table 5). Mean abundances, lengths and biomass, together with the number of counted vs. measured individuals of $D$. sargus and $D$. vulgaris are shown in Appendix, Table 14.

## Effect of habitat

For factor habitat, abundance of $D$. sargus did not show significant differences among different levels (Kruskal-Wallis test, $\mathrm{p}>0.05$; Table 6), despite this was one of the most important species contributing to multivariate differences in abundance between habitats (see chapter 3.1.2). However, significant differences were found both in length and biomass of D. sargus relative to habitat levels (Kruskal-Wallis test, p<0.05; Table 6). Higher average lengths were confirmed in the less complex habitats, hab1 and hab2, when compared to most complex hab4 (Wilcoxon test, $\mathrm{p}<0.05$ ). For biomass, pair-wise comparisons did show that $D$. sargus had greater biomass
in hab1 when compared to other habitat levels. In addition, biomass of $D$. sargus was lower in hab3 than in hab4 (Wilcoxon test, $\mathrm{p}<0.05$; Table 6).

For $D$. vulgaris below MLS, both abundance and biomass were the lowest in hab1 when compared to all other habitats (Wilcoxon test, $\mathrm{p}<0.05$ ) while no habitat effect was found for $D$. vulgaris in general nor $D$. vulgaris above MLS (Wilcoxon test, $\mathrm{p}>0.05$; Table 6).

Table 6: Differences in abundance, length and biomass of key target species between habitats (SBRUV). Results of Kruskal-Wallis test with significant differences ( $\mathrm{p}<0.05$ ) marked with *. Results of pair-wise Wilcoxon test marked with* when inside>outside and ${ }^{* *}$ when inside<outside, only significant results of pair-wise test are shown. MLS: minimum landing size.

| Factor | HABITAT |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Statistic | Main test (Kruskal-Wallis) |  |  | Pair-wise test (Wilcoxon) |  |  |
| Species/ Variable | Abundance | Length | Biomass | Abundance | Length | Biomass |
| D. sargus |  |  |  |  |  |  |
| D. sargus | 0.40 | 0.011* | 0.005* | - | Hab1>Hab4 <br> Hab2>Hab4 | Hab1>Hab2 <br> Hab1>Hab3 <br> Hab1>Hab4 <br> Hab3<Hab4 |
| D. sargus above MLS | 0.70 | - | 0.086 | - | - | - |
| D. sargus below MLS | 0.67 | - | 0.89 | - | - | - |
| D. vulgaris |  |  |  |  |  |  |
| D. vulgaris | 0.104 | 0.66 | 0.43 | - | - | - |
| D. vulgaris above MLS | 0.29 | - | 0.25 | - | - | - |
| D. vulgaris below MLS | 0.035** | - | 0.012** | Hab1<Hab2 <br> Hab1<Hab3 <br> Hab1<Hab4 | - | Hab1<Hab2 <br> Hab1<Hab3 <br> Hab1<Hab4 |

### 3.1.4 Results for the most common non-target species

From the most frequent non-target species, significant differences existed in length of C. julis that was greater inside the reserve than outside (Wilcoxon test, $\mathrm{p}=0.006$; Table 7). For $L$. bergylta, both abundance and biomass were significantly greater inside the reserve than outside (Wilcoxon test, $\mathrm{p}<0.05$; Table 7). For those two species, the observed variables were greater inside the reserve when compared to east outside and/or west outside (Appendix, Table 13). Only in S. cabrilla, the length was lower inside the reserve than in east outside (Wilcoxon test, $\mathrm{p}<0.05$; Appendix, Table 13). The number of counted vs. measured individuals of the three species, mean abundances, length and biomass are shown in Appendix, Table 14.

Table 7: Differences in abundance, length and biomass of the most common non-target species between protection levels and between habitats (SBRUV). Wilcoxon test and Kruskal Wallis test were used to test differences between levels of each factor. Significative p-values ( $\mathrm{p}<0.05$ ) are marked with $*$ when inside>outside and $* *$ when inside<outside.

| Factor | PROTECTION |  |  | HABITAT |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Statistic | Wilcoxon test |  |  | Main test (Kruskal) |  |  |
|  | Abundance | Length (cm) | Biomass (g) | Abundance | Length (cm) | Biomass (g) |
| C. julis | 0.68 | $0.006^{*}$ | 0.13 | 0.46 | 0.057 | 0.85 |
| S. cabrilla | 0.13 | 0.21 | 0.085 | 0.45 | 0.77 | 0.71 |
| L. bergylta | $0.001^{*}$ | 0.063 | $0.015^{*}$ | 0.94 | - | 0.85 |

### 3.2 SDOV

### 3.2.1 Frequency of occurrence, species richness and illegal fishing gear

A total of 15 taxa from 7 different families were sampled with SDOV, with most of the taxa being from Sparidae ( 5 species) and Labridae ( 5 species) families (Appendix, Table 12). Out of all taxa, 8 corresponded to target and 7 to non-target species. Same as for SBRUV, individuals and schools of B. boops appeared in samples but were excluded because of their semi-pelagic lifestyle. From target taxa of Cephalopoda, only S. officinalis appeared within SDOV samples. From target species, D. vulgaris (observed in $82.1 \%$ of the samples) and $D$. sargus ( $74.4 \%$ ) had the highest frequencies of occurrence. From non-target species, C. julis ( $97.4 \%$ ), S. cabrilla ( $46.2 \%$ ) and L. bergylta ( $43.6 \%$ ) were the most frequent (Appendix, Table 12). There were 10 taxa sampled inside and 14 outside the reserve ( 13 in west outside and 12 in east outside). Five species were only found outside (D. cervinus, C. chromis, O. melanura, M. surmuletus, S. officinalis) while one taxa (Mugilidae) appeared only inside the reserve (Appendix, Table 12). No significant differences in species richness per sample were confirmed between inside and outside the reserve (Wilcoxon test, $\mathrm{p}=0.09$; Figure 9), although species richness was significantly greater in west outside than inside (Wilcoxon test, $\mathrm{p}=0.009$ ). Mean richness in samples was equal to $4.22 \pm 0.34$ taxa inside the reserve and $5.19 \pm 0.60$ taxa outside. Illegal fishing gear (octopus traps) was encountered at one occasion inside the reserve (25/05/2018) at Ponta dos Caminhos.


Figure 9: Boxplot of richness by protection (SDOV).

### 3.2.2 Results for community datasets

No statistically significant differences between protection levels were found from multivariate analysis for abundance nor biomass (PERMANOVA, $\mathrm{p}>0.05$; Table 8 ). The only statistically significant difference was found in total abundance of target species below MLS that was greater inside the reserve than outside (Wilcoxon test, $\mathrm{p}=0.04$; Table 8). Total biomass of all community datasets is displayed in Figure 12. Total abundance and total biomass of target species group showed significantly greater mean values in west outside than inside the reserve (Wilcoxon test, $\mathrm{p}<0.05$; Appendix, Table 13). At the same time, target species above MLS and below MLS had significantly greater total abundance and total biomass inside the reserve than in east outside (Wilcoxon test, $\mathrm{p}<0.05$; Appendix, Table 13).

Table 8: Differences in community abundance and biomass between protection levels (SDOV). Results of PERMANOVA and Wilcoxon test. Results of Wilcoxon test with significant p-values ( $\mathrm{p}<0.05$ ) marked with * when inside>outside and ${ }^{* *}$ when inside<outside. MLS: minimum landing size.

| Factor | PROTECTION |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
| Statistic | Main PERMANOVA |  | Wilcoxon test |  |
| Community | Abundance | Biomass | Total abundance | Total biomass |
| All demersal species | 0.17 | 0.06 | 0.30 | 0.20 |
| Target species | 0.10 | 0.24 | 0.57 | 0.53 |
| Target species above MLS | 0.76 | 0.22 | 0.62 | 0.18 |
| Target species below MLS | 0.83 | 0.83 | $0.04^{*}$ | 0.07 |
| Non-target species | 0.27 | 0.35 | 0.21 | 0.06 |



Figure 12: Total biomass by protection (SDOV). Mean $\pm$ standard error displayed. ADS: all demersal species, TS: target species, TS+: target species above MLS, TS-: target species below MLS, NTS: non-target species. Significant differences ( $\mathrm{p}<0.05$ ) marked with *. MLS: minimum landing size.

### 3.2.3 Results for key target species

No significant differences in abundance, length nor biomass were revealed for $D$. sargus for protection (Wilcoxon test, $\mathrm{p}>0.05$; Table 9). For $D$. sargus below MLS, no statistical test was performed as there were only three individuals encountered. Length-frequency distribution of this species is displayed in Figure 13. Individuals of $15-20 \mathrm{~cm}$ class prevailed both outside and inside the reserve. There were no differences in any of $D$. sargus datasets when comparing west outside or east outside to the inside (Wilcoxon test, $\mathrm{p}>0.05$; Appendix, Table 13).

Table 9: Differences in abundance, length and biomass of key target species between protection levels (SDOV). Results of Wilcoxon test with significant p-values ( $\mathrm{p}<0.05$ ) marked with $*$ when inside>outside and $* *$ when inside<outside. MLS: minimum landing size.

| Factor | PROTECTION |  |  |
| :--- | :---: | :---: | :---: |
| Statistic | Wilcoxon test |  |  |
| Species/ Variable | Abundance | Length | Biomass |
| D. sargus |  |  |  |
| D. sargus | 0.91 | 0.96 | 0.76 |
| D. sargus above MLS | 0.73 | - | 0.61 |
| D. sargus below MLS | - | - | - |
| D. vulgaris |  |  |  |
| D. vulgaris | 0.66 | 0.053 | 0.68 |
| D. vulgaris above MLS | 0.91 | - | 0.79 |
| D. vulgaris below MLS | 0.06 | - | $0.04 *$ |



Figure 13: Length-frequency distribution of key target species by protection (SDOV).

There were no significant differences in abundance nor biomass of $D$. vulgaris and $D$. vulgaris above MLS confirmed for protection (Wilcoxon test, $\mathrm{p}>0.05$; Table 9). Differences in length were marginally non-significant, indicating greater length outside than inside the reserve (Wilcoxon test, $\mathrm{p}=0.053$; Table 9, Figure 13). For D. vulgaris below MLS, biomass was significantly greater inside than outside (Wilcoxon test, p<0.04; Table 9, Figure 14) and the same trend was present for abundance of $D$. vulgaris below MLS, although the result was marginally non-significant (Wilcoxon test, $\mathrm{p}=0.06$ ). At the same time, for $D$. vulgaris and $D$. vulgaris above MLS, all variables had significantly greater mean values in west outside than inside the reserve (Appendix, Table 13). The number of counted vs. measured individuals, average abundance, length and biomass of D. sargus and D. vulgaris are shown in Appendix, Table 14.


Figure 14: Biomass of key target species by protection (SDOV). Mean $\pm$ standard error displayed. DS: D. sargus, DSAMLS: D. sargus above MLS, DSBMLS: D. sargus below MLS, DV: D. vulgaris, DV+: D. vulgaris above MLS, DV-: D. vulgaris below MLS. Significant differences ( $\mathrm{p}<0.05$ ) marked with $*$ for inside>outside and $* *$ for inside<outside. MLS: minimum landing size.

### 3.2.4 Results for the most common non-target species

No differences between protection levels were obtained for abundance nor biomass of $C$. julis, S. cabrilla and L. bergylta (Wilcoxon test, p>0.05; Table 10). However, length of C. julis was greater outside than inside the reserve (Wilcoxon test, $\mathrm{p}=0.08$; Table 10), due to greater lengths in east outside when compared to reserve inside (Wilcoxon test, $\mathrm{p}=0.01$; Appendix, Table 13). On the contrary, the abundance of C. julis and of S. cabrilla was significantly greater in west outside than inside (Wilcoxon test, $\mathrm{p}<0.05$; Appendix, Table 13). Due to low number of measured individuals, differences in length of S. cabrilla and L. bergylta were not tested. The numbers of counted vs. measured individuals, mean abundance, length and biomass of the three non-target species are shown in Appendix, Table 14.

Table 10: Differences in abundance, length and biomass of the most common non-target species between protection levels (SDOV). Wilcoxon test and Kruskal Wallis test were used to test differences between levels of each factor. Significative p-values ( $\mathrm{p}<0.05$ ) are marked with $*$ when inside>outside and ${ }^{* *}$ when inside<outside.

| Factor | PROTECTION |  |  |
| :--- | :---: | :---: | :---: |
| Statistic | Wilcoxon test |  |  |
|  | Abundance | Length (cm) | Biomass (g) |
| C. julis | 0.75 | $0.008^{* *}$ | 0.38 |
| S. cabrilla | 0.077 | - | 0.24 |
| L. bergylta | 0.78 | - | 0.74 |

## 4 Methods comparison

### 4.1 Costs comparison

The costs of transportation together with labour costs accounted for the greatest part of the field work costs (Appendix, Table 15). The costs of transportation and field labour were the same for the two methods, as the same number of sampling days and people applied. The costs of bait and diving consumables, which were method specific, represented a relatively negligible part of field sampling costs. Thus, the total value of fieldwork costs per day of sampling was almost equal for the two methods (Appendix, Table 15). However, given that SBRUV provided more samples per day of sampling, the field work costs per sample were $41 \%$ lower for SBRUV than for SDOV (Table 11). Laboratory time was greater for SBRUV than for SDOV (4.6 hours vs. 3.3 hours per sample). This was due to longer time of the SBRUV than SDOV recordings ( 30 mins vs. 4 mins) but also due to other tasks related to video processing (merging of automatically split videos, greater number of calibrations for SBRUV, as three systems were
used, than for SDOV with one system used). This implies greater laboratory labour costs per sample of SBRUV than of SDOV. Anyway, the total variable costs (sum of fieldwork and laboratory costs) of SBRUV per sample remain $26 \%$ below SDOV. When looking at fixed material costs, SDOV represents a more cost demanding option, as it requires acquisition of diving gear on top of stereo-video equipment (Appendix, Table 15).

Table 11: Costs comparison between sampling methods.

|  | Variable costs |  |  |  | Fixed costs |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Fieldwork |  |  | Laboratory |  | Total |
| Sampling method | $\mathrm{N}^{\circ}$ samples <br> per day | Costs <br> per sample | Time <br> per sample | Costs <br> per sample | Costs <br> per sample | Costs <br> per method |
| SBRUV | 10 | $49 €$ | 4.6 h | $27 €$ | $76 €$ | $3000 €$ |
| SDOV | 6 | $83 €$ | 3.3 h | $19 €$ | $102 €$ | $7490 €$ |

### 4.2 Sampling performance comparison

Although part of the SBRUV samples had to be excluded (see reasons of exclusion in 2.2.1), SBRUV still provided greater number of valid samples per day of sampling than SDOV ( 10 vs. 6 samples per day of sampling). Most of the excluded SBRUV samples were successfully resampled, resulting in a total of 60 valid samples out of 63 planned samples. Furthermore, the percentage of measured out of all counted individuals was used as a metric to compare methods efficiency in fish sizes sampling. For the most common species, this percentage was higher for SBRUV than for SDOV (D. sargus: $71 \%>43 \%$, D. vulgaris: $78 \%>56 \%$, C. julis: $53 \%>$ $25 \%$, S. cabrilla: $97 \%>68 \%$ ), only for L. bergylta, SDOV performed better than SBRUV ( $91 \%$ < $100 \%$ ) (Appendix, Table 14).

Total richness in SBRUV samples was double the richness of SDOV samples ( 30 vs. 15 species). Accordingly, species richness per sample was significantly greater in SBRUV than SDOV (Wilcoxon test, $\mathrm{p}<0.001$ ). SDOV sampled successfully only for species that were the most frequent in SBRUV. There were 12 target and 8 non-target species present in SBRUV and absent in SDOV samples (Appendix, Table 12).

The total abundance of all community datasets did not differ significantly between the two methods (Wilcoxon test, $\mathrm{p}>0.05$ ). However, SBRUV sampled for significantly greater total biomass of all demersal species, target species and target species above MLS than SDOV (Wilcoxon test, $\mathrm{p}<0.001$ ). Total biomass of target species below MLS and non-target species did not differ between methods (Wilcoxon test, $\mathrm{p}>0.05$ ). At the same time, total biomass inside
and outside the reserve showed opposite patterns for SBRUV (inside > outside) than for SDOV (inside < outside) for all community datasets, except target species below MLS (Figure 5 and Figure 3.7).

From the most common target and non-target species, D. sargus and S. cabrilla had significantly greater abundance and biomass in SBRUV samples than in SDOV (Wilcoxon test, $\mathrm{p}<0.05)$. For $D$. vulgaris, on the contrary, biomass was greater in SDOV than in SBRUV samples (Wilcoxon test, $\mathrm{p}=0.043$ ) while no difference between methods was found in abundance of this species (Wilcoxon test, $\mathrm{p}=0.085$ ). No differences between methods were obtained for the other two non-target species, C. julis and L. bergylta. Species length did not differ with the method for any of the five tested most common species (Wilcoxon test, $\mathrm{p}>0.05$ ).

## 5 Discussion

The results of our study suggest that positive protection effects exist in Ilhotes do Martinhal marine reserve (Partial Protection Type I), based on SBRUV sampling. Significantly greater mean richness and community biomass were found inside than outside the reserve. The existence of positive protection effect was mainly supported by significantly greater total biomass of target species and target species above minimum landing size (MLS), the two groups being expected to provide the most sensitive response to fishing exclusion, but not of target species below MLS (Harmelin et al., 1995; Mosquera et al., 2000; Micheli et al., 2004; Claudet et al., 2006; Tetreault \& Ambrose, 2007; Molloy et al., 2009; Lester et al., 2009; Horta e Costa et al., 2013a). The fact that length and biomass of D. sargus, as well as biomass of target species above MLS, were significantly greater inside the reserve also when compared to each of the outside areas separately (west outside and east outside), further emphasizes the distinctiveness of the reserve.

Aside from general expectations, the group of non-target species also benefited from protection in terms of their biomass. This result was attributed to L. bergylta, a large-bodied species that accounted for a substantial proportion of biomass of non-target species. In practice, L. bergylta represents a by-catch species with some commercial and subsistence value in recreational fisheries, so it is not completely non-target. In fact, this species has a spawning closure specific for this MPA, revealing fishing interest (Portaria n. ${ }^{\circ}$ 115-A/2011; Portaria n. ${ }^{\circ}$ 115-B/2011). Although by-catch species rarely give a response to protection (Claudet et al., 2010), it could have been envisaged for $L$. bergylta, as the species would not be discarded by fishers due to its greater size (Rakitin \& Kramer, 1996; Mosquera et al., 2010; Malcolm et al., 2018). Conversely,
negative effect of protection was suggested for $S$. cabrilla, a non-target species, same as found in another marine reserve of the Portuguese coast (Horta e Costa et al., 2013a). In this species, the two methods provided complementary results, although significant differences only existed when comparing the reserve inside separately to the west outside and to the east outside.
From the group of target species, $D$. sargus, one of the most commercial species in the region of the MPA (Veiga et al., 2010; Viegas, 2013), showed significantly greater length and biomass inside the reserve. The fact that the abundance did not show significant differences suggests that the significantly higher biomass is mainly driven by the significantly larger sizes of this species inside the reserve. Size and biomass are commonly the first indicators of protection effects, as fisheries generally target larger individuals and increases in fish sizes are a direct response to fishery exclusion (Bianchi et al., 2000; Roberts et al., 2001; Di Franco et al., 2009; Lester et al., 2009; Taylor et al., 2012; Horta e Costa et al., 2013a). In addition, as weight in fish increases exponentially with length, biomass is likely to be the most sensitive metric of early protection effects (Lester et al., 2009). Abundance usually takes longer to answer, being rather an indirect effect of greater sizes, as large females are known to boost the spawning biomass disproportionally (Evans et al., 2008; Barnache et al., 2018). Other factors independent of protection, such as environmental variability, influence the success of recruitment and lead to delayed effects in abundance (Russ \& Alcala, 2004; García-Charton et al., 2004; Shelton \& Mangel, 2011). At community level, the only differences in abundance between the two protection levels were found from multivariate analysis for all demersal species, and for $L$. bergylta at individual species level, but were not confirmed for target species nor target species above MLS. Thus, even after 7 years of protection, positive reserve effects in abundance were still not encountered in most of the cases.

Furthermore, it can be suggested that prohibition of fishing inside of the PPI zone contributes more to a size increase in mature adults of D. sargus and L. bergylta than the temporal closures of artisanal fisheries in those species, which are in place in the buffer zone (Portaria $n^{\circ}{ }^{\circ} 115$ A/2011; Portaria n. ${ }^{\circ}$ 115-B/2011). Response to protection in D. sargus has already been reported from other marine reserves of the Portuguese coast and the Mediterranean even after a few years of protection (two and more years) (Claudet et al., 2006; Horta e Costa et al., 2013a; Belo et al., 2016). Despite its motile character and ability to traverse sandy bottoms at greater distances (several 100s of meters per day) (Vega Fernandéz et al., 2008), D. sargus was found to have limited home ranges, showing site fidelity tendencies, making protection measures efficient when reserve size is adequate (D'Anna et al., 2011; Abecasis et al., 2015; Belo et al., 2016). Such findings, together with our results, suggest that this abundant commercial sparid is
a good candidate for indicator species of protection effects at Ilhotes de Martinhal reserve and other temperate Eastern Atlantic coastal reserves with rocky habitats. The predominance of large individuals of $D$. sargus inside the reserve suggests that this reserve might serve as refuge for spawners, as the sampling period coincided with local spawning season (December to May) (Erzini et al., 2001; Morato et al., 2003). We also found that D. sargus of greater sizes and biomass prevailed in the least structurally complex habitat. Such habitat provides few shelters, thus can only be associated with diel movements related to feeding activity (Morato et al., 2003; Vega Fernandéz et al., 2008).
Furthermore, as both L. bergylta and larger individuals of D. sargus are predators of sea urchins, the effect of protection in those species might result in lower sea urchins' abundance, size and/or biomass and greater corticated macroalgae cover inside the reserve (García-Rubies \& Zabala, 1990; Sala \& Zabala, 1996; Figueiredo et al., 2005; Hereu, 2005; Leitão et al., 2007). Gil Fernandéz et al. (2016) already suggested that effects of trophic cascade were present at Ilhotes do Martinhal reserve, as reflected by lower mean size of sea-urchin Paracentrotus lividus inside the reserve than outside, that correlated with greater turf and foliose algae cover. Those effects were attributed to reserve effect in Diplodus spp. who might control the size of $P$. lividus inside the reserve (Gil Fernandéz et al., 2016).

Contrary to our expectations, for D. vulgaris and D. vulgaris above MLS, another species targeted by commercial fishers and artisanal shore anglers, although less valuable (Gonçalves et al. 2003; Veiga et al., 2010; DGRM, 2018a), negative response to protection was suggested. Except for D. vulgaris below MLS, abundance and biomass was confirmed to be significantly greater outside the reserve, particularly in the west, than inside, as validated by both sampling methods (SBRUV and SDOV). Those results indicate that some level of spatial partitioning might exist between $D$. vulgaris and $D$. sargus, reflecting distinct habitat use due to different food and/or depth preferences in each species (Sala \& Ballestros, 1997), although not investigated in this study. Using physical habitat as explanatory variable, we only found differences in abundance and biomass of $D$. vulgaris below MLS, showing that juveniles of this species avoided the least complex habitat. Those findings are in line with other research that proved juvenile fish to prefer more complex habitats to reduce risk of predation (Sharf et al., 2006). Site fidelity of juveniles of $D$. vulgaris to complex habitats, such as seagrass meadows, has already been documented (Abecasis et al., 2009). However, habitat preferences in juveniles of $D$. vulgaris do not explain the differences in distribution between the two species.

In this context, this study brings new insights about the importance of Ilhotes do Martinhal reserve for D. sargus and D. vulgaris, both commercially important species. Whereas previous
research reported that positive effects in genus Diplodus existed in this reserve (Fernández et al., 2014), we have now demonstrated that $D$. sargus and $D$. vulgaris responded differently. As a result, current limits of Ilhotes do Martinhal reserve do not seem adequate for effective protection of $D$. vulgaris. Given the species strong affinity to west outside locations (Atalaia cliff), an extension of reserve limits over this area offers itself as a solution.

Further habitat effects were found in total abundance of all demersal species and non-target species, showing differences between hab1 and the more complex habitats. Whereas D. sargus had greater abundance in hab1, the other species responsible for the differences had affinity to more complex habitats. Those results are in line with previous findings on habitat preferences in different groups of fish. While small labrid species (C. julis, C. exoletus) were found to require continuous complex habitats, medium-sized sparids ( $D$. sargus) have low requirements on complex habitat connectivity (Vega Fernandéz, 2008). Although statistically nonsignificant, our results suggested that less complex habitats were relatively better represented inside the reserve than more complex habitats. This would theoretically imply lower fish species richness, as greater richness is tied to greater structural habitat complexity (Öhman \& Rajasuriya, 1998; García-Charton \& Pérez-Ruzafa, 2001; Gratwicke \& Speight, 2005), thus working against positive protection effects. Nevertheless, as greater richness was found inside the reserve, physical habitat complexity appears to be suitable enough to ensure protection of demersal species diversity. Apart from that, positive protection effects in $D$. sargus might relate to hab1 relative over-representation inside the reserve, thus eventually confounding the effect of protection itself. Further research is required to provide more information on physical and biological habitat distribution, their quality and use in different species, as well as reserve benefits in terms of biodiversity, based on the entire biological community, including also benthic fish, invertebrate species and algae.
In the absence of data from before reserve implementation, shortcomings of control-effect studies include not only confounding effects of habitat and temporal variability, but also effects of increased fishing pressure in control areas, that might be mistakenly interpreted as enhanced biological productivity inside the reserve (Lester et al., 2009). As illegal octopus traps were repeatedly encountered inside the reserve, further concerns arise regarding fishing intensity inside vs. outside the reserve, opening questions about the true biological carrying capacity of Ilhotes do Martinhal reserve. Supposedly, our results indicate that illegal fishing pressure inside the reserve is less intense than legal fishing pressure outside. However, we do not have information on fishing pressure trends nor biological variables since reserve implementation. Fishing with octopus traps inside the reserve would affect some species from the demersal
community, namely $O$. vulgaris, C. conger and M. helena (Erzini et al., 2008). However, it is possible that other types of illegal fishing gear are also used in the reserve. The presence of illegal fishing in this area reflects the lack of enforcement and possibly the disagreement of locals with this reserve which was top-down implemented (Thaman et al., 2016). At the time of implementation, locals expressed negative perceptions about PNSACV, related to lack of community participation, excessive restrictions, arising conflicts due to recreational fisheries regulation and passive operation (Thaman et al., 2016). Those facts weaken our previous findings about reserve effectiveness and highlight the need for a longterm monitoring program, active management based on local community involvement and legal enforcement being put in place for Ilhotes do Martinhal (Himes, 2003; Mackelworth et al., 2008; Chuenpagdee et al., 2013; Bastista et al., 2015; Gall \& Rodwell, 2016; Thaman et al., 2016).

Compared to SBRUV, no signs of positive protection effects were detected by SDOV. Significantly greater total abundance of target species below MLS and greater biomass of $D$. vulgaris below MLS were found inside the reserve with SDOV. Those results alone are not supportive of protection effects, especially as no positive response was found in target species above MLS. Anyway, the results suggest that juveniles of D. vulgaris might benefit from protection. Conversely, as previously mentioned, D. vulgaris above MLS was found to prefer outside to inside locations. Thus, it can be suggested that this species tends to leave the reserve when adult. The result from SDOV for D. vulgaris below MLS seems complementary to SBRUV which did not unveil differences between the reserve and the outside area.

However, opposite patterns between SDOV and SBRUV are apparent for total biomass, although not significant in SDOV. Contradicting trends in total biomass can be explained by differences in the community sampled between methods. First, individuals of D. sargus above 25 cm , same as some species of greater sizes (D. cervinus, O. melanura, O. vulgaris, D. labrax, $P$. auriga), were rare or absent in SDOV samples. Apart from that, trends in community biomass are partially reflecting trends in biomass of adult $D$. vulgaris, the most frequent species in SDOV, which showed the same trends between methods. In fact, most of the results that compared separately the west outside and the east outside locations to the reserve are compliant for the two methods. For most of the community datasets (except target species above MLS), the west outside locations did not differ from the reserve or even showed significantly greater values of total abundance and total biomass. On the other side, the east outside locations mostly had lower total abundance and total biomass than the reserve. Thus, the findings of both
methods are again supportive of the extention of the current reserve limits to the west outside locations.

Differences between SDOV and SBRUV results are related to lower statistical power of SDOV data, as an outcome of lower richness and lower number of individuals measured, on one side, and limited spatial coverage of the SDOV sampling design applied, on the other side. Significantly lower richness and biomass of SDOV samples likely resulted from negative behavioural response in fish to divers (Francour et al., 1999; Pais \& Cabral, 2017; Pais \& Cabral, 2018) in combination with low visibility conditions (Tessier et al., 2013). Whereas in SBRUV, fish tends to approach the cameras while attracted by the bait, making low visibility less restrictive, in SDOV, most of the fish swims away from the divers. The only species that was more efficiently sampled with SDOV than SBRUV was D. vulgaris, showing greater biomass from SDOV. This supports that $D$. vulgaris, especially larger individuals, are not very shy of divers (Kulbicki, 1998; Guidetti et al., 2008). As observed by Kulbicki (1998) and Watson \& Harvey (2007), in some species, large individuals tend to come closer to divers than smaller individuals. The 'tolerance' to divers in $D$. vulgaris might also be related to the fact that this species, unlike D. sargus, is not preferentially targeted by spearfishers in Portugal (Assis et al., 2018). Furthermore, D. vulgaris is easily identifiable, based on the two-banded pattern, which favoured its sampling using SDOV, contrary to other species that are small and/or have cryptic patterns, such as C. julis, Symphodus spp. or S. cabrilla (Tessier et al., 2013). In this study, opposite results were found for individual species, as in the case of length of $C$. julis. We believe that in SDOV, small individuals of this species are being systematically undersampled as they often appear too far from the cameras to be measured. This might cause higher average lengths when less individuals are measured, potentially generating biased results.

Further, SDOV studies might opt for cameras with more sensitive sensors, as those could improve the sampling of cryptic fish under limiting visibility conditions. Apart from that, a wider common FOV of cameras might increase the number of individuals measured, being another suggested improvement to SDOV performance. As another limitation, a narrower sampling design applied to SDOV than SBRUV ( 39 vs. 60 samples). This was a result of the same number of days at sea allocated to both methods while SDOV provided less samples per day than SBRUV. Taking into consideration that marine organisms follow heterogeneous, patchy spatial distribution (García-Charton \& Pérez-Ruzafa, 2001), greater spatial coverage of sampling should correct for spatial variability due to this heterogeneity. SBRUV samples, separated by a minimum distance of 250 m , addressed better this spatial heterogeneity issue, providing representative mean values for the sampled locations. Although SDOV, with 10 m
distance between transects, generated reasonable replicates for a concrete 'patch', the number of sites sampled would need to be increased to guarantee greater representativeness of mean values for the entire sampled location. Thus, we believe that increased number of sampling sites would make the SDOV method more competitive in terms of statistical power of the biological data.

Furthermore, the area effectively sampled by SBRUV was not estimated, due to the unknown area of bait attraction for fish (Taylor et al., 2013; Haggitt et al., 2014). As a result, the compared sampling units differed between methods (ex. MaxN from 30 min stationary stereo-video vs. continuous count from 30 mx 6.2 m swimmable transect), same as in other comparative studies (Watson et al., 2005; Langlois et al., 2010). This raises questions about direct comparability of values obtained using two different sampling units, especially when knowing that total abundance and biomass will increase in SDOV as a function of transect length. Alternatively, MaxN should also be used in SDOV to avoid the influence of transect length.

Overall, SBRUV was confirmed to be a more efficient monitoring tool than SDOV, not only in terms of biological indicators but also in terms of cost-efficiency. As field-sampling costs per sample are low, they compensate for additional laboratory post-processing costs, same as concluded by Watson et al. $(2005 ; 2010)$ and Langlois et al. (2010). A comparison with classical UVC should be done for demersal fish and commercial invertebrate community, as UVC is expected to sample for more fish species than both SBRUV (Langlois et al., 2006; Colton \& Swearer, 2010) and SDOV (Greene \& Alevizon, 1989; Pelletier et al., 2011; Holmes et al., 2013; Tessier et al., 2013). UVC also allows to sample for benthic fish, thus providing data on protection effects in the whole fish community associated with rocky reefs (Monteiro et al., 2012; Henriques et al., 2013). Although an underestimation of richness and density of cryptobenthic species occurs in UVC, the method still seems to be more efficient in sampling of this group of species than video-methods (Willis, 2001; Watson et al., 2005; Lowry et al., 2012; Wilson et al., 2018). On top of behavioural responses in fish to divers, UVC also carries observers bias, such as variability between observers in community sampled and lower accuracy of fish length measurements when compared to the stereo-video (Harvey et al., 2002; Williams et al., 2006). UVC represents a more competitive option than SDOV in terms of costs, as it does not involve laboratory post-processing of videos (Holmes et al., 2013; Tessier et al., 2013). However, same is valid for UVC as for SDOV in terms of sampling design spatial coverage which should be representative enough for the area under question. This would imply more days at sea than for SBRUV and influence UVC cost competitiveness.

## Conclusions

It can be concluded that, after 7 years of protection, our results in richness and biomass of demersal fish and commercial invertebrates support the existence of eventual protection effects at Ilhotes do Martinhal marine reserve. The groups usually most sensitive to protection (target species and, particularly, target species above MLS) were found to thrive inside the reserve in terms of their biomass, with D. sargus being a candidate indicative species of protection effects. It was confirmed that this marine reserve is also beneficial for large by-catch species ( $L$. bergylta). At the same time, some other species, both commercial (D. vulgaris) and noncommercial (S. cabrilla), showed affinity to locations which are situated outside of the marine reserve. Influence of other habitat characteristics than physical habitat should be investigated to fully distinguish protection effect from other spatial preferences. An ongoing monitoring of the marine reserve is needed to validate our findings and to accompany further trends in abundance which was not affirmative of protection effects. Although responses in abundance to protection might be delayed due to environmental factors, the interference with illegal fishing might be another explanation of non-persuasive results in abundance. This is the reason why active management, based on local community participation and reserve enforcement, should become a priority if this marine reserve is aimed to be successful.

When to methods comparison, SBRUV represents a suitable sampling tool for long-term monitoring of MPAs such as Ilhotes do Martinhal marine reserve, as this method was able to detect differences indicative of protection effects in both demersal fish community and in individual species. At the same time, SBRUV provided overall greater sampling performance and cost effectiveness than SDOV. Our results showed that SDOV sampled for narrower species community and undersampled small and cryptic species, most probably due to negative diver effect and visibility limitations, when opposed to effect of bait. We suggest that comparison with classical UVC method should be done for Ilhotes do Martinhal marine reserve to decide on the most appropriate non-destructive monitoring method to be used.

The findings of this study are especially relevant for the management of Ilhotes do Martinhal marine reserve, as they will serve as a basis for reassessment of protection measures in place, including potential adjustments in zonation, and for the development of a long-term monitoring scheme.

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## 7 Appendix

Table 12: Target and non-target fish and invertebrate taxa by sampling method. Commercial status (CS): target (T) or non-target (NT), frequency of occurrence (FO) in \%, presence $(\checkmark)$ or absence (X) of the taxa IN: inside and/or OUT: outside the reserve, not sampled at all with the method $(-)$.

|  |  |  | SBRUV |  |  | SDOV |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Family | Scientific Name | CS | $\begin{aligned} & \text { FO } \\ & (\%) \end{aligned}$ | IN | OUT | $\begin{aligned} & \mathrm{FO} \\ & (\%) \end{aligned}$ | IN | OUT |
| Congridae | Conger conger | T | 1.7 | X | $\checkmark$ | - | - | - |
| Muraenidae | Muraena helena | T | 3.3 | $\checkmark$ | $\checkmark$ | - | - | - |
| Gadidae | Trisopterus luscus | T | 1.7 | X | $\checkmark$ | - | - | - |
| Carangidae | Pseudocaranx dentex | NT | 1.7 | X | $\checkmark$ | - | - | - |
|  | Trachurus trachurus | T | 8.3 | $\checkmark$ | $\checkmark$ | - | - | - |
| Labridae | Centrolabrus exoletus | NT | 30.0 | $\checkmark$ | $\checkmark$ | 15.4 | $\checkmark$ | $\checkmark$ |
|  | Coris julis | NT | 98.3 | $\checkmark$ | $\checkmark$ | 97.4 | $\checkmark$ | $\checkmark$ |
|  | Ctenolabrus rupestris | NT | 41.7 | $\checkmark$ | $\checkmark$ | 23.1 | $\checkmark$ | $\checkmark$ |
|  | Labrus bergylta | NT | 36.7 | $\checkmark$ | $\checkmark$ | 43.6 | $\checkmark$ | $\checkmark$ |
|  | Labrus mixtus | NT | 3.3 | $\checkmark$ | $\checkmark$ | - | - | - |
|  | Symphodus spp. | NT | 18.3 | $\checkmark$ | $\checkmark$ | 25.6 | $\checkmark$ | $\checkmark$ |
| Haemulidae | Plectorhinchus mediterraneus | NT | 1.7 | $\checkmark$ | X | - | - | - |
| Moronidae | Dicentrarchus labrax | T | 10.0 | $\checkmark$ | $\checkmark$ | - | - | - |
| Mugilidae | Mugilidae | T | 30.0 | $\checkmark$ | $\checkmark$ | 25.6 | $\checkmark$ | $\checkmark$ |
| Mullidae | Mullus surmuletus | T | 1.7 | X | $\checkmark$ | 5.1 | X | $\checkmark$ |
| Pomacentridae | Chromis chromis | NT | 3.3 | X | $\checkmark$ | 10.3 | X | $\checkmark$ |
| Serranidae | Serranus atricauda | NT | 3.3 | X | $\checkmark$ | - | - | - |
|  | Serranus cabrilla | NT | 81.7 | $\checkmark$ | $\checkmark$ | 46.2 | $\checkmark$ | $\checkmark$ |
| Sparidae | Diplodus cervinus | T | 20.0 | $\checkmark$ | $\checkmark$ | 2.6 | X | $\checkmark$ |
|  | Diplodus sargus | T | 96.7 | $\checkmark$ | $\checkmark$ | 74.4 | $\checkmark$ | $\checkmark$ |
|  | Diplodus vulgaris | T | 95.0 | $\checkmark$ | $\checkmark$ | 82.1 | $\checkmark$ | $\checkmark$ |
|  | Oblada melanura | T | 20.0 | $\checkmark$ | $\checkmark$ | 10.3 | X | $\checkmark$ |
|  | Pagrus auriga | T | 10.0 | $\checkmark$ | X | - | - | - |
|  | Pagrus pagrus | T | 1.7 | $\checkmark$ | X | - | - | - |
|  | Sarpa salpa | T | 11.7 | $\checkmark$ | $\checkmark$ | - | - | - |
|  | Sparus aurata | T | 3.3 | X | $\checkmark$ | - | - | - |
|  | Spondyliosoma cantharus | T | 6.7 | $\checkmark$ | $\checkmark$ | - | - | - |
| Loliginidae | Loligo vulgaris | T | 5.0 | $\checkmark$ | X | - | - | - |
| Octopodidae | Octopus vulgaris | T | 20.0 | $\checkmark$ | $\checkmark$ | - | - | - |
| Sepiidae | Sepia officinalis | T | 5.0 | X | $\checkmark$ | 5.1 | X | $\checkmark$ |

Table 13: Comparison of results for protection between methods. Protection effect: Yes (Y) marked * when inside $>$ outside and ${ }^{* *}$ when inside < outside, No (N). For comparison between reserve inside (I) and west outside (W), and reserve inside (I) and east outside (E), only significant results are shown ( $\mathrm{p}<0.05$ ). Results are from Wilcoxon test.

|  | BRUV |  | SDOV |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Protection effect | W vs. I vs. E | Protection effect | W vs. I vs. E |
|  | Richness |  |  |  |
| Richness per sample | Y* | I > E | N | W > I |
|  | Community assemblages |  |  |  |
|  | Abundance |  |  |  |
| All demersal species | N | W > I > E | N | no diff |
| Target species | N | $\mathrm{W}>\mathrm{I}>\mathrm{E}$ | N | W > I |
| Target species above MLS | N | no diff | N | I > E |
| Target species below MLS | N | no diff | Y* | $\mathrm{I}>\mathrm{E}$ |
| Non-target species | N | I > E | N | no diff |
|  | Biomass |  |  |  |
| All demersal species | Y* | $\mathrm{I}>\mathrm{E}$ | N | W > I |
| Target species | Y* | $I>E$ | N | $\mathrm{W}>\mathrm{I}>\mathrm{E}$ |
| Target species above MLS | Y* | $\mathrm{W}<\mathrm{I}>\mathrm{E}$ | N | $\mathrm{I}>\mathrm{E}$ |
| Target species below MLS | N | $\mathrm{I}>\mathrm{E}$ | N | $\mathrm{I}>\mathrm{E}$ |
| Non-target species | Y* | I > E | N | no diff |
|  | Key-target species |  |  |  |
|  | Abundance |  |  |  |
| D. sargus | N | $\mathrm{I}>\mathrm{E}$ | N | No diff |
| D. sargus above MLS | N | $\mathrm{I}>\mathrm{E}$ | N | No diff |
| D. sargus below MLS | N | no diff | - | - |
| D. vulgaris | N | W > I | N | W > $>$ > E |
| D. vulgaris above MLS | $\mathrm{Y}^{* *}$ | $\mathrm{W}>\mathrm{I}$ | N | $\mathrm{W}>\mathrm{I}>\mathrm{E}$ |
| D. vulgaris below MLS | N | $\mathrm{W}>\mathrm{I}$ | N | $\mathrm{I}>\mathrm{E}$ |
|  | Length |  |  |  |
| D. sargus | Y* | W < I | N | No diff |
| D. vulgaris | N | No diff | N | W > I |
|  | Biomass |  |  |  |
| D. sargus | Y* | W < I > E | N | No diff |
| D. sargus above MLS | Y* | $\mathrm{W}<\mathrm{I}>\mathrm{E}$ | N | No diff |
| D. sargus below MLS | N | no diff | N | No diff |
| D. vulgaris | $\mathrm{Y}^{* *}$ | W > I | N | W > $>$ > |
| D. vulgaris above MLS | Y** | $\mathrm{W}>\mathrm{I}$ | N | $\mathrm{W}>\mathrm{I}>\mathrm{E}$ |
| D. vulgaris below MLS | N | No diff | Y* | $\mathrm{I}>\mathrm{E}$ |
|  | Non-target species |  |  |  |
|  | Abundance |  |  |  |
|  | Protection | W vs. I. vs. E | Protection | W vs. I. vs. E |
| C. julis | N | I > E | N | W > I |
| S. cabrilla | N | No diff | N | $\mathrm{W}>\mathrm{I}$ |
| L. bergylta | Y* | W < I > E | N | No diff |
|  | Length |  |  |  |
| C. julis | Y* | W < I | $\mathrm{Y}^{* *}$ | I < E |
| S. cabrilla | N | No diff | - | - |
| L. bergylta | N | - | - | - |
|  | Biomass |  |  |  |
| C. julis | N | I > E | N | No diff |
| S. cabrilla | N | $\mathrm{I}<\mathrm{E}$ | N | No diff |
| L. bergylta | Y* | $\mathrm{W}<\mathrm{I}$ | N | No diff |

Table 14: Abundance, length, biomass and $\%$ of measured individuals of the most common species. Mean $\pm$ standard error shown for abundance, length and biomass. Abundance and biomass are per sample, length is mean individual length.

| variable | SBRUV |  |  | SDOV |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | whole area | inside | outside | whole area | inside | outside |
|  | D. sargus |  |  |  |  |  |
| abundance | $5.3 \pm 0.5$ | $5.8 \pm 0.8$ | $4.9 \pm 07$ | $4.6 \pm 0.9$ | $4.3 \pm 1.3$ | $4.8 \pm 1.3$ |
| length (cm) | $19.5 \pm 0.3$ | $20.2 \pm 0.45$ | $18.8 \pm 0.32$ | $18.9 \pm 0.40$ | $18.8 \pm 0.57$ | $19.0 \pm 0.6$ |
| biomass (g) | $823.0 \pm 103.8$ | $1079.7 \pm 189.6$ | $594.4 \pm 97.6$ | $502.6 \pm 134.8$ | $437.6 \pm 151.8$ | $558.4 \pm 216.9$ |
| $\mathbf{N}^{\circ}$ counted | 315 | 150 | 165 | 178 | 77 | 101 |
| $\mathbf{N}^{\circ}$ measured | 224 | 113 | 111 | 77 | 36 | 41 |
| \% measured | 71 | 75 | 67 | 43 | 47 | 41 |
|  | D. vulgaris |  |  |  |  |  |
| abundance | $5.0 \pm 0.5$ | $4.7 \pm 0.9$ | $5.1 \pm 0.6$ | $7.4 \pm 1.1$ | $7.3 \pm 1.4$ | $7.5 \pm 1.6$ |
| length (cm) | $14.7 \pm 0.3$ | $14.7 \pm 0.4$ | $14.7 \pm 0.3$ | $15.4 \pm 0.3$ | $14.8 \pm 0.4$ | $16.0 \pm 0.5$ |
| biomass (g) | $280.9 \pm 36.6$ | $242.0 \pm 63.3$ | $310.7 \pm 43.1$ | $554.1 \pm 111.1$ | $465.9 \pm 113.3$ | $629.7 \pm 183.3$ |
| $\mathrm{N}^{\circ}$ counted | 301 | 122 | 179 | 290 | 132 | 158 |
| $\mathbf{N}^{\circ}$ measured | 236 | 86 | 148 | 162 | 80 | 81 |
| \% measured | 78 | 71 | 83 | 56 | 61 | 52 |
|  | C. julis |  |  |  |  |  |
| abundance | $14.4 \pm 1.8$ | $14.5 \pm 2.6$ | $14.4 \pm 2.5$ | $20.1 \pm 3.2$ | $15.9 \pm 2.5$ | $23.7 \pm 5.5$ |
| length (cm) | $11.0 \pm 0.2$ | $11.5 \pm 0.3$ | $10.6 \pm 0.2$ | $10.8 \pm 0.3$ | $10.3 \pm 0.3$ | $11.6 \pm 0.4$ |
| biomass (g) | $235.5 \pm 26.3$ | $276.5 \pm 45.3$ | $204.1 \pm 30.4$ | $371.7 \pm 75.1$ | $239.1 \pm 46.6$ | $485.4 \pm 130.0$ |
| $\mathbf{N}^{\circ}$ sampled | 867 | 377 | 490 | 784 | 286 | 498 |
| $\mathbf{N}^{\circ}$ measured | 455 | 189 | 266 | 201 | 117 | 83 |
| \% measured | 53 | 50 | 54 | 25 | 41 | 17 |


|  | S. cabrilla |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| abundance | $1.45 \pm 0.1$ | $1.6 \pm 0.2$ | $1.3 \pm 0.2$ | $0.7 \pm 0.1$ | $0.5 \pm 0.2$ | $0.9 \pm 0.2$ |  |  |
| length (cm) | $13.5 \pm 0.3$ | $13.6 \pm 0.5$ | $13.4 \pm 0.3$ | $13.3 \pm 0.8$ | $12.3 \pm 0.9$ | $14.7 \pm 1.2$ |  |  |
| biomass (g) | $55.5 \pm 6.8$ | $64.9 \pm 10.6$ | $48.4 \pm 8.75$ | $21.3 \pm 6.3$ | $11.7 \pm 5.7$ | $29.5 \pm 10.4$ |  |  |
| $\mathbf{N}^{\circ}$ counted | 87 | 42 | 45 | 28 | 9 | 19 |  |  |
| $\mathbf{N}^{\circ}$ measured | 84 | 41 | 43 | 19 | 6 | 13 |  |  |
| \% measured | 97 | 98 | 96 | 68 | 67 | 68 |  |  |
|  | L. bergylta |  |  |  |  |  |  |  |
| abundance | $0.38 \pm 0.07$ | $0.62 \pm 0.1$ | $0.21 \pm 0.08$ | $0.62 \pm 0.2$ | $0.5 \pm 0.2$ | $0.71 \pm 0.2$ |  |  |
| length (cm) | $25.95 \pm 1.5$ | $27.41 \pm 1.9$ | $23.84 \pm 2.3$ | $25.65 \pm 1.7$ | $26.7 \pm 3.0$ | $24.7 \pm 1.9$ |  |  |
| biomass (g) | $153.2 \pm 44.6$ | $196.66 \pm 63.5$ | $120.9 \pm 62.0$ | $119.1 \pm 46.07$ | $110.0 \pm 74.8$ | $126.9 \pm 58.4$ |  |  |
| $\mathbf{N}^{\circ}$ counted | 23 | 16 | 8 | 17 | 8 | 9 |  |  |
| $\mathbf{N}^{\circ}$ measured | 21 | 13 | 8 | 17 | 8 | 9 |  |  |
| \% measured | 91 | 81 | 100 | 100 | 100 | 100 |  |  |

Table 15: Costs of field work, laboratory work and fixed material costs of sampling methods.

|  | SBRUV | SDOV |
| :---: | :---: | :---: |
| Total field work costs (per day of sampling) | 490 | 500 |
| Transportation |  |  |
| Total boat <br> Boat fuel (average consumption) per day <br> Boat rental or usage per day <br> Total car <br> Car fuel (Faro-Sagres-Faro) <br> Car road fees (Faro-Sagres-Faro) |  |  |
| Fieldwork labour costs |  |  |
| man-day costs 3x (skipper 1x, sampling technicians 2x) |  |  |
| Fieldwork consumables | 10 | 24 |
| Total diving consumables tank fillings (air) per day ( 4 x ) diving insurance per day ( 4 x ) <br> Total baited cameras consumables <br> Bait (per day) | n/a <br> n/a <br> n/a <br> 10 <br> 10 | 24 <br> 16 <br> 8 <br> n/a <br> $\mathrm{n} / \mathrm{a}$ |
| Laboratory costs (per day of work) |  |  |
| Laboratory labour costs |  |  |
| man-day costs 1x |  |  |
| Fixed costs (all material) | 3000 | 7490 |
| Stereo-video system | 3000 | 1790 |
| Cameras equipment <br> Action cameras (SBRUV: SK4 4x, GoPro Hero3 2x; SDOV: SK4 2x) <br> - SK4 (SBRUV 4x, SDOV 2x) <br> - GoPro Hero3 (SBRUV 2x) <br> Memory cards (SBRUV: 6x, SDOV: 2x) <br> Extra batteries (SBRUV: 6x) <br> Stereo-video hardware <br> SBRUV frame 3 x <br> Calibration cube <br> Stereo-video analysis and data storage <br> iMac for VidSync <br> External storage disk | $\begin{gathered} \hline \mathbf{7 5 0} \\ 600 \\ 200 \\ 400 \\ 100 \\ 50 \\ \mathbf{6 5 0} \\ 600 \\ 50 \\ \mathbf{1 6 0 0} \\ 1500 \\ 100 \end{gathered}$ | $\begin{gathered} \hline \mathbf{1 4 0} \\ 100 \\ 100 \\ - \\ 40 \\ \mathrm{n} / \mathrm{a} \\ \mathbf{5 0} \\ \mathrm{n} / \mathrm{a} \\ 50 \\ \mathbf{1 6 0 0} \\ 1500 \\ 100 \end{gathered}$ |
| Diving material | n/a | 5700 |
| personal diving equipment 2 x <br> diving tanks 8 x <br> Oxygen kit | n/a <br> n/a <br> n/a | $\begin{gathered} \hline 3600 \\ 1600 \\ 500 \end{gathered}$ |

Table 16: SIMPER results for community abundance and biomass based on protection (SBRUV). The table shows average abundance and average biomass in protection level (inside, outside), average dissimilarity value, dissimilarity to standard deviation ratio (diss/SD), \% contribution of each species to dissimilarity and cumulative \% contribution to dissimilarity.

|  | PROTECTION |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Average abundance |  | Dissimilarity |  |  |  |
| Species | inside | outside | avg diss | diss/SD | contrib\% | cum.\% |
| All demersal species |  |  | 43.25 |  |  |  |
| C. julis | 3.51 | 3.36 | 7.54 | 1.24 | 17.44 | 17.44 |
| D. vulgaris | 1.90 | 2.15 | 4.35 | 1.34 | 10.05 | 27.49 |
| D. sargus | 2.24 | 2.03 | 4.31 | 1.21 | 9.96 | 37.45 |
| S. cabrilla | 1.14 | 0.99 | 2.76 | 1.03 | 6.38 | 43.83 |
| Mugilidae | 0.21 | 0.54 | 2.50 | 0.84 | 5.77 | 49.60 |
| L. bergylta | 0.62 | 0.19 | 2.45 | 1.11 | 5.68 | 55.28 |
| S. salpa | 0.12 | 0.60 | 2.34 | 0.35 | 5.42 | 60.70 |
| C. rupestris | 0.55 | 0.32 | 2.24 | 0.98 | 5.18 | 65.87 |
| O. melanura | 0.40 | 0.19 | 2.14 | 0.67 | 4.94 | 70.81 |
| C. exoletus | 0.35 | 0.32 | 1.88 | 0.84 | 4.34 | 75.15 |
| D. cervinus | 0.35 | 0.09 | 1.54 | 0.74 | 3.56 | 78.71 |
| O. vulgaris | 0.27 | 0.15 | 1.45 | 0.68 | 3.35 | 82.06 |
| Symphodus spp. | 0.22 | 0.19 | 1.35 | 0.65 | 3.11 | 85.17 |
| D. labrax | 0.19 | 0.10 | 1.01 | 0.43 | 2.34 | 87.51 |
| P. auriga | 0.25 | 0.00 | 0.98 | 0.53 | 2.27 | 89.78 |
| T. trachurus | 0.12 | 0.06 | 0.68 | 0.43 | 1.58 | 91.36 |
| Target species |  |  | 45.61 |  |  |  |
| D. sargus | 2.24 | 2.03 | 8.60 | 1.19 | 18.85 | 18.85 |
| D. vulgaris | 1.90 | 2.15 | 8.58 | 1.32 | 18.81 | 37.66 |
| Mugilidae | 0.21 | 0.54 | 4.83 | 0.84 | 10.59 | 48.26 |
| O. melanura | 0.40 | 0.19 | 4.02 | 0.68 | 8.81 | 57.07 |
| S. salpa | 0.12 | 0.60 | 3.76 | 0.37 | 8.24 | 65.31 |
| O. vulgaris | 0.27 | 0.15 | 2.96 | 0.67 | 6.49 | 71.80 |
| D. cervinus | 0.35 | 0.09 | 2.90 | 0.73 | 6.36 | 78.15 |
| D. labrax | 0.19 | 0.10 | 1.99 | 0.44 | 4.37 | 82.52 |
| P. auriga | 0.25 | 0.00 | 1.95 | 0.53 | 4.27 | 86.80 |
| T. trachurus | 0.12 | 0.06 | 1.17 | 0.42 | 2.56 | 89.35 |
| L. vulgaris | 0.13 | 0.00 | 1.00 | 0.34 | 2.19 | 91.55 |
| Target species above MLS |  |  | 45.09 |  |  |  |
| D. vulgaris | 0.91 | 1.34 | 11.55 | 1.22 | 25.61 | 25.61 |
| D. sargus | 1.86 | 1.70 | 10.72 | 1.07 | 23.77 | 49.38 |
| Mugilidae | 0.19 | 0.33 | 5.05 | 0.72 | 11.21 | 60.59 |
| D. cervinus | 0.35 | 0.06 | 4.60 | 0.72 | 10.20 | 70.79 |
| D. labrax | 0.17 | 0.06 | 2.62 | 0.43 | 5.80 | 76.60 |
| S. salpa | 0.12 | 0.15 | 2.49 | 0.41 | 5.51 | 82.11 |
| T. trachurus | 0.12 | 0.06 | 1.89 | 0.43 | 4.19 | 86.30 |
| P. auriga | 0.17 | 0.00 | 1.77 | 0.42 | 3.92 | 90.22 |
| Target species below MLS |  |  | 42.31 |  |  |  |


| D. vulgaris | 1.31 | 1.66 | 18.36 | 1.17 | 43.39 | 43.39 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| D. sargus | 0.27 | 0.25 | 8.14 | 0.65 | 19.23 | 62.62 |
| O. vulgaris | 0.18 | 0.12 | 6.62 | 0.53 | 15.64 | 78.26 |
| S. cabrilla | 0.11 | 0.08 | 3.30 | 0.40 | 7.81 | 86.07 |
| S. salpa | 0.00 | 0.13 | 1.60 | 0.20 | 3.78 | 89.85 |
| P. pagrus | 0.06 | 0.00 | 1.32 | 0.21 | 3.12 | 92.97 |
| Non-target species |  |  | 39.20 |  |  |  |
| C. julis | 3.51 | 3.46 | 15 | 1.27 | 38.25 | 38.25 |
| S. cabrilla | 1.14 | 1.02 | 5.93 | 0.94 | 15.13 | 53.38 |
| L. bergylta | 0.62 | 0.19 | 5.17 | 1.01 | 13.18 | 66.56 |
| C. rupestris | 0.55 | 0.33 | 4.52 | 0.96 | 11.53 | 78.09 |
| C. exoletus | 0.35 | 0.33 | 3.69 | 0.86 | 9.4 | 87.49 |
| Symphodus spp. | 0.22 | 0.19 | 2.7 | 0.65 | 6.89 | 94.38 |
|  | Average biomass |  | Dissimilarity |  |  |  |
| Species | inside | outside | avg diss | diss/SD | contrib\% | cum.\% |
| All demersal species |  |  | 56.94 |  |  |  |
| D. sargus | 30.67 | 20.98 | 8.12 | 1.14 | 14.26 | 14.26 |
| L. bergylta | 9.28 | 4.41 | 4.94 | 0.88 | 8.67 | 22.93 |
| D. cervinus | 11.12 | 1.09 | 4.91 | 0.67 | 8.62 | 31.56 |
| Mugilidae | 4.86 | 7.64 | 4.7 | 0.71 | 8.25 | 39.81 |
| D. vulgaris | 13.05 | 15.97 | 4.35 | 1.38 | 7.64 | 47.44 |
| C. julis | 15.03 | 12.69 | 4.00 | 1.24 | 7.03 | 54.47 |
| O. vulgaris | 5.69 | 3.20 | 3.71 | 0.63 | 6.51 | 60.98 |
| S. salpa | 2.11 | 6.49 | 3.25 | 0.37 | 5.71 | 66.69 |
| D. labrax | 5.97 | 2.16 | 3.1 | 0.44 | 5.45 | 72.14 |
| O. melanura | 4.00 | 1.34 | 2.28 | 0.68 | 4.00 | 76.14 |
| S. cabrilla | 7.03 | 5.80 | 2.18 | 1.20 | 3.83 | 79.97 |
| P. auriga | 4.39 | 0.00 | 1.73 | 0.43 | 3.04 | 83.01 |
| L. vulgaris | 3.33 | 0.00 | 1.46 | 0.28 | 2.56 | 85.57 |
| Symphodus spp. | 2.03 | 1.44 | 1.28 | 0.63 | 2.25 | 87.82 |
| C. rupestris | 2.06 | 1.42 | 1.14 | 1.05 | 2.01 | 89.82 |
| T. trachurus | 1.61 | 0.72 | 1.05 | 0.38 | 1.85 | 91.68 |
| Target species |  |  | 59.15 |  |  |  |
| D. sargus | 30.67 | 20.98 | 11.95 | 1.10 | 20.20 | 20.20 |
| D. cervinus | 11.12 | 1.09 | 6.82 | 0.68 | 11.52 | 31.73 |
| Mugilidae | 4.86 | 7.64 | 6.65 | 0.71 | 11.24 | 42.96 |
| D. vulgaris | 13.05 | 15.97 | 6.31 | 1.34 | 10.67 | 53.63 |
| O. vulgaris | 5.69 | 3.20 | 5.36 | 0.61 | 9.06 | 62.69 |
| D. labrax | 5.97 | 2.16 | 4.25 | 0.45 | 7.19 | 69.89 |
| S. salpa | 2.11 | 6.49 | 4.19 | 0.39 | 7.08 | 76.97 |
| O. melanura | 4.00 | 1.34 | 3.25 | 0.67 | 5.50 | 82.46 |
| P. auriga | 4.39 | 0.00 | 2.47 | 0.43 | 4.18 | 86.64 |
| L. vulgaris | 3.33 | 0.00 | 1.81 | 0.28 | 3.05 | 89.7 |
| S. officinalis | 0.00 | 2.23 | 1.34 | 0.24 | 2.27 | 91.97 |
| Target species above MLS |  |  | 55.33 |  |  |  |
| D. sargus | 26.51 | 20.43 | 11.06 | 1.01 | 19.98 | 19.98 |


| D. vulgaris | 8.76 | 13.43 | 8.58 | 1.25 | 15.50 | 35.48 |
| :--- | ---: | ---: | ---: | ---: | ---: | ---: |
| D. cervinus | 11.12 | 1.20 | 8.55 | 0.69 | 15.46 | 50.94 |
| Mugilidae | 4.47 | 6.40 | 7.05 | 0.70 | 12.75 | 63.69 |
| D. labrax | 5.43 | 1.57 | 4.77 | 0.44 | 8.63 | 72.32 |
| S. salpa | 1.96 | 2.70 | 3.03 | 0.40 | 5.47 | 77.79 |
| P. auriga | 4.04 | 0.00 | 2.58 | 0.39 | 4.67 | 82.46 |
| O. vulgaris | 2.57 | 1.36 | 2.43 | 0.34 | 4.39 | 86.84 |
| L. vulgaris | 3.33 | 0.00 | 2.27 | 0.28 | 4.10 | 90.94 |
| Target species below MLS |  |  | 49.51 |  |  |  |
| D. vulgaris | 7.62 | 8.86 | 15.91 | 1.36 | 32.14 | 32.14 |
| O. vulgaris | 3.69 | 2.57 | 13.65 | 0.56 | 27.57 | 59.71 |
| D. sargus | 1.97 | 1.71 | 8.84 | 0.62 | 17.86 | 77.57 |
| S. cantharus | 1.28 | 0.75 | 4.90 | 0.39 | 9.90 | 87.47 |
| D. labrax | 0.00 | 0.95 | 1.67 | 0.20 | 3.38 | 90.85 |
| Non-target species |  |  | 47.61 |  |  |  |
| L. bergylta | 9.28 | 4.68 | 14.19 | 0.98 | 29.81 | 29.81 |
| C. julis | 15.03 | 13.48 | 13.51 | 1.21 | 28.38 | 58.18 |
| S. cabrilla | 7.03 | 6.16 | 7.90 | 1.03 | 16.59 | 74.77 |
| Symphodus spp. | 2.03 | 1.53 | 3.97 | 0.64 | 8.33 | 83.1 |
| C. rupestris | 2.06 | 1.51 | 3.88 | 0.98 | 8.15 | 91.25 |

Table 17: SIMPER results for community abundance based on habitats (SBRUV). The table shows average abundance and average biomass in habitats (hab1, hab2, hab3, hab4), average dissimilarity value, dissimilarity to standard deviation ratio (diss/SD), \% contribution of each species to dissimilarity and cumulative $\%$ contribution to dissimilarity.

|  | HABITAT |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Average a | undance |  | Dissim | ilarity |  |
| Hab1 vs. Hab2 | Hab1 | Hab2 | avg diss | diss/SD | contrib\% | cum.\% |
| All demersal species |  |  | 44.82 |  |  |  |
| C. julis | 2.96 | 3.32 | 7.47 | 1.20 | 16.66 | 16.66 |
| D. sargus | 2.36 | 2.03 | 4.21 | 1.06 | 9.40 | 26.06 |
| D. vulgaris | 1.55 | 1.98 | 4.09 | 1.31 | 9.13 | 35.19 |
| S. cabrilla | 0.85 | 1.26 | 3.39 | 1.08 | 7.56 | 42.75 |
| S. salpa | 0.13 | 0.77 | 2.94 | 0.34 | 6.55 | 49.30 |
| Mugilidae | 0.43 | 0.33 | 2.64 | 0.75 | 5.89 | 55.19 |
| C. rupestris | 0.00 | 0.53 | 2.22 | 1.03 | 4.96 | 60.16 |
| L. bergylta | 0.35 | 0.36 | 2.07 | 0.87 | 4.61 | 64.77 |
| O. melanura | 0.24 | 0.40 | 1.98 | 0.80 | 4.42 | 69.19 |
| Symphodus spp. | 0.14 | 0.36 | 1.78 | 0.76 | 3.98 | 73.17 |
| D. cervinus | 0.29 | 0.20 | 1.62 | 0.73 | 3.61 | 76.78 |
| C. exoletus | 0.12 | 0.33 | 1.51 | 0.75 | 3.36 | 80.14 |
| D. labrax | 0.08 | 0.30 | 1.42 | 0.49 | 3.17 | 83.31 |
| O. vulgaris | 0.29 | 0.07 | 1.38 | 0.66 | 3.08 | 86.39 |
| P. auriga | 0.18 | 0.13 | 1.11 | 0.58 | 2.48 | 88.87 |
| T. trachurus | 0.18 | 0.07 | 0.95 | 0.52 | 2.13 | 90.99 |


| Hab1 vs. Hab3 | Hab1 | Hab3 | avg diss | diss/SD | contrib\% | cum.\% |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| All demersal species |  |  | 43.86 |  |  |  |
| C. julis | 2.96 | 3.86 | 8.77 | 1.19 | 19.99 | 19.99 |
| D. sargus | 2.36 | 1.94 | 5.52 | 1.35 | 12.58 | 32.57 |
| D. vulgaris | 1.55 | 2.28 | 4.87 | 1.32 | 11.11 | 43.68 |
| S. cabrilla | 0.85 | 1.04 | 3.29 | 1.09 | 7.49 | 51.17 |
| C. rupestris | 0.00 | 0.58 | 2.77 | 1.04 | 6.32 | 57.49 |
| Mugilidae | 0.43 | 0.44 | 2.71 | 0.88 | 6.18 | 63.67 |
| L. bergylta | 0.35 | 0.45 | 2.21 | 0.93 | 5.04 | 68.71 |
| O. vulgaris | 0.29 | 0.18 | 1.75 | 0.73 | 3.99 | 72.70 |
| D. cervinus | 0.29 | 0.18 | 1.66 | 0.72 | 3.79 | 76.49 |
| C. exoletus | 0.12 | 0.25 | 1.37 | 0.54 | 3.12 | 79.61 |
| S. salpa | 0.13 | 0.18 | 1.11 | 0.52 | 2.54 | 82.15 |
| O. melanura | 0.24 | 0.00 | 1.05 | 0.54 | 2.40 | 84.54 |
| Symphodus spp. | 0.14 | 0.09 | 0.9 | 0.47 | 2.05 | 86.6 |
| T. trachurus | 0.18 | 0.00 | 0.79 | 0.46 | 1.81 | 88.4 |
| P. auriga | 0.18 | 0.00 | 0.76 | 0.45 | 1.73 | 90.14 |
| Hab1 vs. Hab4 | Hab1 | Hab4 | avg diss | diss/SD | contrib\% | cum.\% |
| All demersal species |  |  | 46.38 |  |  |  |
| C. julis | 2.96 | 3.69 | 8.76 | 1.33 | 18.88 | 18.88 |
| D. vulgaris | 1.55 | 2.45 | 5.37 | 1.39 | 11.57 | 30.45 |
| D. sargus | 2.36 | 2.07 | 3.91 | 1.37 | 8.42 | 38.88 |
| S. cabrilla | 0.85 | 1.10 | 3.05 | 1.17 | 6.57 | 45.44 |
| C. rupestris | 0.00 | 0.65 | 2.58 | 1.26 | 5.57 | 51.01 |
| Mugilidae | 0.43 | 0.40 | 2.56 | 0.87 | 5.52 | 56.53 |
| O. melanura | 0.24 | 0.41 | 2.36 | 0.60 | 5.10 | 61.63 |
| C. exoletus | 0.12 | 0.60 | 2.34 | 0.98 | 5.04 | 66.66 |
| S. salpa | 0.13 | 0.45 | 2.11 | 0.33 | 4.56 | 71.22 |
| L. bergylta | 0.35 | 0.35 | 1.89 | 0.87 | 4.08 | 75.30 |
| O. vulgaris | 0.29 | 0.24 | 1.63 | 0.77 | 3.52 | 78.82 |
| D. cervinus | 0.29 | 0.12 | 1.42 | 0.68 | 3.07 | 81.88 |
| Symphodus spp. | 0.14 | 0.20 | 1.14 | 0.57 | 2.47 | 84.35 |
| P. auriga | 0.18 | 0.08 | 0.92 | 0.52 | 1.98 | 86.33 |
| T. trachurus | 0.18 | 0.06 | 0.90 | 0.51 | 1.94 | 88.28 |
| L. vulgaris | 0.14 | 0.06 | 0.85 | 0.42 | 1.82 | 90.10 |

