The Seabed Makes the Dolphins: Physiographic Features Shape the Size and Structure of the Bottlenose Dolphin Geographical Units

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Abstract: The common bottlenose dolphin ( Tursiops truncatus ) is a cosmopolitan delphinid, regularly present in the Mediterranean Sea. According to previous studies, this dolphin tends to form resident geographical units scattered on the continental shelf. We investigated how the physiographic characteristics of the area of residence, with special reference to the size and shape of the continental shelf, affect the home range and the group size of the local units. We analysed and compared data collected between 2004–2016 by 15 research groups operating in different study areas of the Mediterranean Sea: the Alboran Sea, in the South-Western Mediterranean, the Gulf of Lion and the Pelagos Sanctuary for the marine mammals, in the North-Western Mediterranean, and the Gulf of Ambracia, in the North-Central Mediterranean Sea. We have found that in areas characterised by a wide continental platform, dolphins have wider home ranges and aggregate into larger groups. In

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areas characterized by a narrow continental platform, dolphins show much smaller home ranges and aggregate into smaller groups. The results obtained from this collective research effort highlight the importance of data sharing to improve our scientific knowledge in the field of cetaceans and beyond.

**Keywords:** bottlenose dolphin; Mediterranean Sea; habitat; home range; group size

1. Introduction

The common bottlenose dolphin—*Tursiops truncatus* Montagu, 1821—hereafter referred to as bottlenose dolphin, is a cosmopolitan Delphinidae that can be found worldwide in all the oceans. Its distribution is usually confined between 45°12′ parallels, in tropical and temperate waters of both hemispheres, except for the North Atlantic, where it can reach the 65°12′ parallel [1–5].

This wide distribution of the species is associated with a certain level of morphometric differentiation among the different areas and latitudes, which seems consistent with the presence of a globally distributed offshore ecotype and several inshore ecotypes, scattered along the coasts [6–20].

The inshore bottlenose dolphin communities show a high level of site fidelity, with home range sizes that can vary from tens of km² [21] to hundreds of km² [22]. According to a global synthesis provided by Wells and Scott [3], the home range of the inshore bottlenose dolphins generally includes areas of about 100–150 km², even if wider home ranges have been reported for local geographical units [23]. It has been hypothesized that the home range sizes and movement patterns of this species depend on habitat differences [24], food resources [25] and reproductive resources [26].

In the context of the Mediterranean Sea, the bottlenose dolphin is considered a regularly present species and the second most sighted cetacean after the striped dolphin, *Stenella coeruleoalba* [27–31].

According to Notarbartolo di Sciara and Demma [29], the Mediterranean bottlenose dolphin would be more closely related to the Atlantic inshore ecotype, while Cañadas et al. [32,33], who studied this species in the context of the Alboran Sea, suggested a closer link with the offshore Atlantic ecotype. This apparent contradiction could in fact derive from the different ecological habits of the dolphins that inhabit the Alboran Sea (as also emerges from the present study). Gaspari et al. [34] found that samples from deeper parts of the Mediterranean (e.g., Ionian Sea) were genetically more similar to samples from the Atlantic offshore ecotype and suggested a potential differentiation between offshore and inshore ecotypes also in the Mediterranean context. However, the results from TursioMed, a networking project compiling census data at the Mediterranean level and involving 29 institutes from eight Mediterranean countries, covering 13 years of survey effort from 2004 to 2016 [35], showed that most sightings of the species occurred within the 200 m isobath marking the border of the continental shelf, while sightings outside this limit have been reported to be quite rare despite the sampling effort. This pattern of distribution seems quite consistent in all sampled study areas, with the already mentioned exception of the Alboran Sea, possibly confirming the inshore habit of the Mediterranean bottlenose dolphin, as originally stated by Notarbartolo Sciara and Demma [29]. Therefore, it is unclear to what extent the genetic differences found by Gaspari and co-authors, who examined mainly stranded animals, should be correlated to ecological habits (see also [14]).

According to Moura et al. [36], the Mediterranean and Black Sea (inshore) populations derive from a globally distributed offshore population, which would explain the presence of an “offshore” genetic inheritance in Mediterranean bottlenose dolphins, regardless of their current ecological habits.

Most studies based on photo identification data are consistent in describing the Mediterranean bottlenose dolphin as a primarily resident species. According to Gnone et al. [37], who studied the movement patterns of this species in the Pelagos Sanctuary (a
specially protected area in the northwest of the Mediterranean basin, see below), the bottlenose dolphins move around their usual area of residence within a maximum distance of 50 kms. The resident behaviour and a local specialization, especially in feeding techniques, seem to produce a segregation between neighbouring dolphins and a clustering of the (meta) population in geographical and demographical units [37]. According to Carnabuci et al. [38], the connectivity (or disconnectivity) through the bottlenose dolphin units living in the Pelagos Sanctuary retraces the landscape features and its habitat breakages (see also [39]). The dolphins living in the Tuscany Archipelagos, for example, and those inhabiting the waters off the northwest coast of Corsica, are separated by a remarkable habitat breakage, delimited by the Corsica’s “finger” (Cape Corso). The east side (Tuscany Archipelagos) is characterized by shallow waters and sandy/muddy ecosystems, while the west side (northwest coast of Corsica) presents a typical narrow rocky platform and a steep slope. The “ecological distance” seems to overcome the geographical distance, producing a clear separation of the two units despite the geographical proximity.

The specialization on the residence habitat and the isolation between geographical units could be responsible for the genetic structure described by Natoli et al. [40] and, on a finer scale, by Gaspari et al. [34] and Brotons et al. [41].

As bottlenose dolphins form resident units scattered over the continental shelf, it is crucial to understand how the physiographic characteristics of the area of residence, which can be very different even in neighbouring areas, can affect the geographical units, shaping their structure.

For this purpose, we analysed six geographical units of bottlenose dolphins living in different areas of the Mediterranean Sea, comparing their home ranges (both as individuals and as a whole unit), their aggregation behaviour (group size), their abundance and density.

2. Material and Methods

2.1. Study Areas

The data were collected by different research groups in four areas of the Mediterranean basin, recognized as Important Marine Mammal Areas (IMMA) or Specially Protected Area of Mediterranean Importance (SFAMI), and in the neighbouring waters (Figure 1).
2.1.1. Alboran Sea

The Alboran Sea is the westernmost area of the Mediterranean. Located at the entrance of the basin, it is connected to the Atlantic Ocean through the Strait of Gibraltar and represents a transition area from the ocean to the semi-enclosed Mediterranean Sea. It covers a water area of approximately 56,000 km$^2$, between Spain and Morocco, with an average depth of 445 m and a maximum depth of 1500 m in its eastern portion. The seabed shows a complex profile and is crossed, in its central part, by an important chain of seamounts, the Alboran Ridge. The area has been recognized as IMMA, where the bottlenose dolphin is reported as a qualifying species [42]. Most of the data analysed in the present study come from the northern side of the Alboran Sea.

2.1.2. Gulf of Lion

The Gulf of Lion is a wide embayment, located in the easternmost portion of the French Mediterranean coast, up to the border with Spain. It covers an area of approximately 13,000 km$^2$ over the continental shelf, with an average depth of about 250 m and no relevant habitat breakages. The Gulf of Lion is also included in the IMMAs list for the bottlenose dolphin identified as a qualifying species [43].

2.1.3. Pelagos Sanctuary

The Pelagos Sanctuary for Mediterranean Marine Mammals is the largest Mediterranean Marine Protected Area and it is listed as the biggest SPAMI within the framework of the Barcelona Convention. It is located in the north-western portion of the basin, and it covers an area of about 87,500 km$^2$, extending over waters belonging to the territorial waters of France, Italy and the Principality of Monaco (and the adjacent international waters), including Corsican and north Sardinian coasts [44]. It comprises pelagic areas, deep water zones and the continental shelf, with remarkable oceanographic and physio-
graphic differences. The western portion is characterized by typical rocky coasts, with a narrow continental shelf, furrowed by several submarine canyons, and a wide bathyal plain (2500–2700 m) [45]; the eastern area, on the contrary, has a much wider continental shelf, with typical sandy/muddy ecosystems. Compared to other Mediterranean regions, this area is characterized by very high levels of offshore primary productivity, induced by the interaction between geomorphologic, oceanographic and climatic factors [46]. All cetaceans regularly observed in the Mediterranean Sea can be found in this region, including the common bottlenose dolphin [37,39,47–52]. The Sanctuary area partially overlaps with the “North Western Mediterranean Sea, Slope and Canyon System IMMA”, which includes the bottlenose dolphin as (secondary) qualifying species [53].

2.1.4. Gulf of Ambracia

The Gulf of Ambracia (or Gulf of Amvrakikos) is a semi-enclosed gulf in north-western Greece. It has a total surface of about 400 km$^2$, and it is connected to the Ionian Sea through a 700 m wide canal. The average depth is only 30 m, the maximum depth is 63 m, and its bottom mostly consists of mud or sand. Its waters are rich in nutrients and characterized by abundant wildlife. The dolphins of the Gulf of Ambracia are genetically differentiated from the surrounding Mediterranean metapopulation [54]. The area is recognized as IMMA, where the bottlenose dolphin is the only qualifying species [55].

2.2. Data Collection

We analysed the data collected between 2004–2016 by 15 different research groups operating in the above described Mediterranean regions (see Table 1) and shared on the Intercet platform (www.intercet.it, accessed on 10 May 2022) within the TursioMed project [35].
Table 1. Summary of information on the research groups involved in the study.

| Research Group                                              | Interchet Code | Study Area                                                                 | Sampling Period |
|--------------------------------------------------------------|----------------|----------------------------------------------------------------------------|-----------------|
| Alnilam Research and Conservation                            | ARC            | Alboran Sea (Spain)                                                        | 2004–2011       |
| SUBMON                                                       | SBN            | Catalonia (Spain)                                                          | 2010–2015       |
| Association BREACH                                          | BR             | Gulf of Lion (France)                                                      | 2006–2016       |
| EcoOcean Institut and partners *                             | EOI            | Gulf of Lion, French Riviera, western Corsica (France)                     | 2005–2015       |
| Tethys Research Institute—CSR                               | TRI            | Western Liguria (Italy), French Riviera (France)                          | 2004–2016       |
| Fondazione Acquario di Genova                                | DM             | Eastern Liguria (Italy)                                                    | 2004–2016       |
| University of Genoa—DISTAV                                  | UGE            | Eastern Liguria, Tuscany (Italy)                                           | 2005–2008       |
| CE.TU.S Cetacean Research Centre                            | CC             | Tuscany (Italy)                                                            | 2004–2016       |
| ARPAT                                                        | AT             | Tuscany (Italy)                                                            | 2010–2011       |
| Pelagos                                                      | PE             | Tuscany (Italy)                                                            | 2011            |
| Association CARI                                            | CA             | Northern Corsica (France)                                                  | 2013–2015       |
| GECEM (MIRACETI)                                            | GC             | Gulf of Lion, French Riviera, Corsica (France)                            | 2004–2016       |
| Office de l’Environnement de la Corse                        | BB             | Strait of Bonifacio (France, Italy)                                        | 2009–2012       |
| Bottlenose Dolphin Research Institute                       | BDR            | North eastern Sardinia (Italy)                                             | 2004–2013       |
| Tethys Research Institute—IDP                               | IDP            | Gulf of Ambracia (Greece)                                                  | 2004–2016       |

*Dedicated surveys were carried out by expert researchers between March–October, in favourable weather conditions (sea state < 4 in the Douglas scale), following random tracks within predefined sampling areas. Each research team used a GPS device to record the effort tracks (the routes travelled by the research vessel during the sampling activity) and the sighting points of the target species. Researchers counted the number of individuals observed in each group and collected photographic data for individual photo-identification [56].

2.3. Data Analysis

2.3.1. Distribution

The GIS system ArcGIS 9.3 was used to analyse the georeferenced data (the effort tracks and the sighting points of the target species) collected by the research groups involved in the study.

2.3.2. Photographic Data and Matching Process

The research groups selected their photographic data separately, but following common rules: (a) the quality of the image; and (b) the distinctiveness of the animals [35].

(a) Photographic quality was based on focus, lighting (the sun should be behind the camera for proper dorsal fin illumination), angle of the fin to the photographer (the fin
should be in an orthogonal position with respect to the camera, to avoid deformation of the profile), contrast between dorsal fin and background and visibility of the fin [57–59].

(b) Identification value was based on the presence of notches, deformities, unusual fin shapes, scars and discoloration [60]. Notches, deformities and unusual fin shapes were considered to be permanent marks [58] and were used as primary distinctive elements for the photo-identification of the animals. Scars and discolorations were only used in association with primary distinctive elements.

The catalogues curated by the single research groups were then pooled to be matched. The matching between catalogues was performed with aid of FinfindR (https://github.com/haimeh/finFindR, accessed on 10 May 2022), a software for computer-assisted recognition and identification of the bottlenose dolphin dorsal fins [61]. Each recapture had to be confirmed visually by an experienced researcher, and if in doubt, a second person was asked to confirm the match.

We used the Simple Ratio Index (SRI) [62] to measure the degree of sharing between catalogues, calculated according to the following formula:

\[
SRI = \frac{X}{(a + b - X)}
\]  

(a = total number of individuals in catalogue a; b = total number of individuals in catalogue b; and X = number of individuals shared between a and b).

The SRI can vary from 0 (no individuals shared between the catalogues matched) to 1 (all the individuals in the catalogue a are present in the catalogue b and vice versa), depending on the reciprocal size of the two catalogues and the number of individuals shared.

2.3.3. Connectivity Analysis and Cluster Identification

Following the matching, a network outline was drawn using the Spring embedding visualization [63,64] in NetDraw [65]. This was performed selecting those individuals sighted and recognised through photographic identification on at least 4 occasions (including the first identification event), 4 being the average number of times each individual was sighted, as in Carnabuci et al. [38].

A more severe selection could have provided more info about the “intimate” structure within each cluster, but we were actually looking to the network “macro-structure” on a Mediterranean scale. Furthermore, we could verify that the size of the minimum convex polygon (see below) did not increase with the increase in the number of captures >4, reassuring us this was an appropriate choice.

The connectivity analysis was then performed on a selection of 704 individuals of the 2604 photo-identified (about 27%). Two animals were assumed to be associated in the network “macro-structure” on a Mediterranean scale. Furthermore, we could verify that the size of the minimum convex polygon (see below) did not increase with the increase in the number of captures >4, reassuring us this was an appropriate choice.

The connectivity analysis was then performed on a selection of 704 individuals of the 2604 photo-identified (about 27%). Two animals were assumed to be associated in the network if sighted in the same group on at least one occasion, following a 0–1 criterion (0, never sighted together; 1, sighted together at least once).

To identify the clusters within the network, we carried out a Girvan–Newman analysis, based on edge betweenness measurements [66–69]. The best division for the network was identified using a modularity index Q, where the highest Q value indicates the best division [69,70]. This index varies between 0 (community structure no better than in a random network) and 1 (strong community structure) and it can be considered meaningful if it falls in the range 0.3–0.7 [71].

Each identified cluster was assigned a different colour, which was maintained in all the figures to facilitate the comprehension of the analytical process.

2.3.4. Mapping the Different Geographical Units

Following the connectivity analysis and cluster identification, all the sightings of the individuals selected (all the individuals with at least 4 captures, see above) were then plotted on the map using the same cluster colours, in order to identify the areas of occupancy of the bottlenose dolphins belonging to each cluster.
2.3.5. Movement Analysis through Minimum Convex Polygon (MCP)

For each of the selected individuals, we have drawn the minimum convex polygon [72] as a simplified representation of its home range. Each (convex) polygon includes all the sighting points of the same individual and describes its maximum recorded movements. Since the distribution of sightings clearly shows that the bottlenose dolphins are confined within the border of the continental shelf area (except for the bottlenose dolphins inhabiting the Alboran Sea), we cropped the MCPs following the 200 m isobath profile. This crop has been implemented for all the individual MCPs, with the exception of the dolphins inhabiting the Alboran Sea (see above) and the ones residing in the Gulf of Ambracia (which is entirely included within the continental shelf).

Cropping the MCP along the 200 m isobath is a needed correction to avoid serious bias when drawing the MCPs. The bias derives from the curvature of the coast profile and is proportional to its degree. Since the shelf usually follows the coastal contour, an MCP drawn across a concave coastline will result in the inclusion of pelagic waters outside the 200 isobath that were actually free of sightings and thus will result in an overestimated area.

A boxplot was used for a descriptive analysis of MCP area values and to identify outliers within each cluster. We then compared the average size of the individual MCPs belonging to each cluster, excluding the outliers identified through the boxplot analysis. The statistical significance of the differences in the average size of the cluster MCPs was assessed with a Kruskal–Wallis test for independent samples (a non-parametric test was chosen in consideration of the Kolmogorov–Smirnov normality test results). For the pairwise comparison of the clusters, the significance was adjusted by the Bonferroni correction for multiple tests.

The average MCP of the different clusters were then compared to the corresponding MP and to the average width of the continental platform in the area of occupancy, to test for possible correlation through a nonlinear regression analysis.

To identify the home range of the different clusters/geographical units, we overlapped the MCPs of the single individuals belonging to the same cluster, so as to obtain a complex multi-polygon (MP).

The home range of a geographical unit can be underestimated if the sampling effort was concentrated in an area too small to describe the movements of the dolphins in their full extent. We then calculated, for each cluster, the ratio between the average MCP (the average of the individual MCPs) and the MP of the cluster, to obtain an “overlap index”. The overlap index can be used as a measure of the extent of effort relative to the size of the average MCP. For example, if the overlap index approaches 1, it means that the total sampled area approximates the average MCP, suggesting a possible bias given by effort coverage (since the MCP cannot be larger than the same sampled area). Therefore, the overlap index can provide information on the reliability of the results obtained.

2.3.6. Group Size

For each cluster/geographical unit, we calculated the average group size, considering all the sightings where at least one of the selected individuals was present. We then compared the mean group sizes across clusters/geographical units, testing the statistical significance of the differences through a Kruskal–Wallis test for independent samples (also in this case a non-parametric test was chosen in consideration of the Kolmogorov–Smirnov normality test results). For the pairwise comparison of the clusters, the significance was adjusted by the Bonferroni correction for multiple tests.

We used a nonlinear regression analysis to test the relationship between: the individual’s MCP and the average size of the groups in which the same individual was sighted, defined as the “individual group size” (IGS), and the average MCP and the average group size in the different clusters. The statistical significance of the correlations found was verified with the ANOVA Regression test.
2.3.7. Abundance Estimates and Density

We estimated the abundance of each cluster/geographical unit using a closed population mark-recapture model [73], performed with Capture [74,75], within the Mark program [76]. Through this analysis, we produced single year abundance estimates of the animals identified by means of permanent natural markings, considering all the sightings where at least one individual of the above-described selection (all the individuals with at least 4 captures) was identified. This model, applied to the annual dataset, was considered the most appropriate approach according to the above home range analysis and literature consultation [37,58,59,77].

Mark–recapture models for closed populations rely on a series of assumptions: (a) the mark (or the recognition system) should be reliable during the study period; (b) the capture of an animal should not modify the probability of the same individual being captured again; (c) the population should be considered closed, meaning that emigration/immigration and birth/death events should not affect the size of the population during the study period, neither should the portion of marked and unmarked animals change within the population; (d) all of the individuals of the population should have the same probability of being captured at each sampling event. Strong violations of these assumptions may lead to bias in the abundance estimate and, therefore, a proper validation of the data collected should be done in advance [78].

In our study, assumption (a) is respected, as we considered permanent marks for individual photo-identification. Although these may change over time (new marks may be added), the use of multiple elements reduces the risk for false negative or positive matches, especially if the estimate is performed over a single year period. Assumption (b) is also respected, as the photographic capture technique does not significantly affect the behavioural response of the individual to the next capture event. Regarding the assumption (c), the bottlenose dolphins are long living and slow breeding animals; according to Rossi et al. [79], the annual fertility rate in the Mediterranean context is approximately 0.29–0.41 and is compensated by a similar death rate. In relation to immigration and emigration events, we based the assumption of closure on the previous cluster analysis. In fact, the connections between clusters are few over the entire research period (13 years) and are minimal (if any are present) within the years selected for the abundance estimates. This is consistent with the resident behaviour exhibited by the dolphins in the study areas, in agreement with Gnone et al. [37]. Assumption (d) is usually the most difficult to satisfy [58,80] and its violation may lead to an underestimation of the population size. Within this study, capture probabilities are unlikely to be constant, owing to the multiple research groups conducting the data collection. For this reason, a time dependent and/or heterogeneity model, which allows a certain tolerance of assumptions (d) should be used. We chose the time dependent variant (Chao Mt) of the Chao model [81] (Chao et al., 1992), based on Wilson et al. [58], Chilvers and Corkeron [60] and Gnone et al. [37].

For each cluster, we chose the annual estimate with the maximum number of individuals captured (photo-identified), considering it as the most reliable (best annual estimate). Finally, the population density was calculated as the ratio between the abundance estimate and the size of the cluster MP.

3. Results

3.1. Distribution

A total of 306,388 kms of sampling effort was performed by the research groups in their respective study areas (Table 2, Figure 2), and 3927 sightings of bottlenose dolphins were recorded (Table 2, Figure 3). In the surveyed areas, the distribution of bottlenose dolphins was clearly confined within the 200 m isobath that marks the boundary of the continental shelf, except for the Alboran Sea, where dolphins have been sighted even in deeper waters, confirming previous findings from Cañadas et al. [32].
Table 2. The sampling effort implemented by each research group (Effort (km)), the resulting sightings of bottlenose dolphins (Sightings Tt), the sightings during which photographic data for photo-identification of the individuals were collected (Sight. Photo-ID) and the total of individuals photo-identified (Ind. Photo-ID).

| Research Group (Interet Code) | Sampling Period | Effort (km) | Sightings Tt | Sight. Photo-ID | Ind. Photo-ID |
|-------------------------------|-----------------|-------------|--------------|----------------|---------------|
| ARC                           | 2004–2011       | 24,200      | 210          | 100            | 539           |
| SBN                           | 2010–2015       | 692         | 10           | 10             | 52            |
| BR                            | 2006–2016       | 3754        | 95           | 75             | 494           |
| EOI                           | 2005–2015       | 48,758      | 65           | 38             | 281           |
| TRI                           | 2004–2016       | 66,770      | 32           | 27             | 121           |
| DM                            | 2004–2016       | 26,382      | 233          | 210            | 263           |
| UGE                           | 2005–2008       | 15,591      | 49           | 38             | 169           |
| CC                            | 2004–2016       | 18,710      | 432          | 330            | 321           |
| AT                            | 2010–2011       | 2864        | 27           | 27             | 126           |
| PE                            | 2011            | 2842        | 34           | 27             | 199           |
| CA                            | 2013–2015       | 1359        | 6            | 6              | 17            |
| GC                            | 2004–2016       | 28,542      | 213          | 190            | 665           |
| BB                            | 2009–2012       | 3494        | 46           | 41             | 95            |
| BDR                           | 2004–2013       | 12,328      | 1637         | 683            | 48            |
| IDP                           | 2004–2016       | 50,102      | 838          | 205            | 123           |
| **TOTAL**                     |                 | **306,388** | **3927**     | **2007**       | **3513**      |

Figure 2. Sampling tracks implemented by each research group involved in the different study areas (see also Tables 1 and 2). The dark line marks border of the continental shelf (200 m isobath) in all the maps.
3.2. Photographic Data and Matching Process

In 2007 sightings (out of a total of 3927), the researchers collected photographic data for the photo-identification of individuals, and a gross total of 3513 dolphins were photo-identified by the different groups (Table 2, Figure 4). The comparison between catalogues resulted in 909 positive matches, lowering the net total to 2604 distinctively identified individuals (Table 3). Only in 13 cases (highlighted orange in Table 3) did the Simple Ratio Index (SRI) exceed 0.1, always between research groups sampling in the same area or in neighbouring areas, confirming the strong site-fidelity of the bottlenose dolphins.
Table 3. The result of the matching between photo-ID catalogues (the number of individuals shared across the different catalogues) and relative SRI between catalogues (in parentheses). The SRI can vary from 0 (no individuals shared between catalogues) to 1 (all the individuals in the first catalogue are present in the second one and vice versa). In green the cells with SRI > 0, in orange the cells with SRI > 0.1.

| Code | Total | ARC | SBN | BR | EOI | TRI | DM | UGE | CC | AT | PE | CA | GC | BB | BDR | IDP |
|------|-------|-----|-----|----|-----|-----|----|-----|----|----|----|----|----|----|-----|-----|
| ARC  | 539   | 539 |     |    |     |     |    |     |    |    |    |    |    |    |     |     |
| SBN  | 52    | 0   | 52  |    |     |     |    |     |    |    |    |    |    |    |     |     |
| BR   | 494   |     | 18  | 494|     |     |    |     |    |    |    |    |    |    |     |     |
| EOI  | 281   | 2   | 11  | 117| 281 |     |    |     |    |    |    |    |    |    |     |     |
| TRI  | 121   | 3   | 12  | 8  | 121 |     |    |     |    |    |    |    |    |    |     |     |
| DM   | 263   |     | 40  | 95 | 169 |     |    |     |    |    |    |    |    |    |     |     |
| UGE  | 169   |     |     | 36 | 183 | 90  | 321|     |    |    |    |    |    |    |     |     |
| CC   | 321   |     |     | 36 | 183 | 90  | 321|     |    |    |    |    |    |    |     |     |
| AT   | 126   | 4   | 35  | 23 | 38  | 126 | 72  |     |    |    |    |    |    |    |     |     |
| PE   | 199   |     |     | 17 | 86  | 39  | 94  | 39  | 199 |     |    |    |    |    |     |     |
| CA   | 17    |     |     | 3  | 6   | 0   | 0   | 0   |    | 17  |     |    |    |    |     |     |
| GC   | 665   | 2   | 23  | 151| 127 | 655 | 7  | 0   | 3  | 3  | 10 | 95 |    |    |     |     |
| BB   | 95    |     |     | 0  | 0   | 0   | 0   | 0   | 0  | 0  | 0  | 0  | 10 | 95 |     |     |
| BDR  | 48    |     |     | 0  | 0   | 0   | 0   | 0   | 0  | 0  | 0  | 0  | 2  | 16 | 48  |     |
| IDP  | 123   |     |     | 0  | 0   | 0   | 0   | 0   | 0  | 0  | 0  | 0  | 0  | 0  | 123 |     |
| Total (gross) | 3513 | 539 | 52  | 494| 281 | 121 | 263 | 169 | 321 | 126 | 199 | 17 | 655 | 95  | 48  |
| Positive matches | 909  |     |     |    |     |     |    |     |    |    |    |    |    |    |     |     |
| Total (net)     | 2604 |     |     |    |     |     |    |     |    |    |    |    |    |    |     |     |

3.3. Connectivity Analysis and Cluster Identification

The connectivity analysis was performed on the photo-identified dolphins with at least 4 “captures”, for a total of 703 individuals. The Spring embedding visualization shows five major aggregations around the points of densest connection (Figure 5) and the Girvan–Newman analysis identifies six clusters within the same network (best division for the network was identified with the value of Qmax = 0.656), based on the betweenness of the bonds (Figure 5): (A) composed by 95 individuals; (B) composed by 132 individuals; (C) composed by 261 individuals; (D) composed by 66 individuals; (E) composed by 36 individuals; and (F) composed by 113 individuals.
3.4. Mapping the Different Geographical Units

Following the connectivity analysis and cluster identification, all the sightings of the selected individuals were plotted on the map, using the same cluster colours coding as in Figure 4. The clusters clearly correspond to geographical units with their own area of distribution (Figure 6).

Figure 5. The connectivity network of the photo-identified bottlenose dolphins with at least four captures (703 individuals) according to the Spring embedding lay out. A Girvan–Newman analysis was performed to identify the clusters within the network according to the betweenness of the bonds (see the different colours).

Figure 6. The sighting positions of the photo-identified bottlenose dolphins with at least four captures (703 individuals). The colours correspond to the belonging cluster (see Figure 5).
3.5. Movement Analysis through Minimum Convex Polygon (MCP)

For each of the 703 individuals selected, we have drawn the respective Minimum Convex Polygon (MCP), as shown in Figure 7. When calculating the surface area (in km$^2$) of the individual MCPs, the portion lying outside the 200 m isobath, delimiting the boundaries of the bottlenose dolphin habitat, was cropped (cropping has not been applied to the MCPs of the dolphins inhabiting the Alboran Sea, whose habitat does not appear to be so sharply confined to the continental shelf, see above).

![Figure 7. The minimum convex polygons (MCPs) of the photo-identified bottlenose dolphins with at least four captures (703 individuals). The colours correspond to the belonging cluster (see Figure 4). When calculating the surface area (in km$^2$) of the individual MCPs, the portion lying outside the 200 m isobath, delimiting the boundaries of the bottlenose dolphin habitat, was cropped, except for cluster A (purple colour), residing in the Alboran Sea (see the main text for further details).](image)

The boxplot analysis identified a total of 19 outliers among the clusters (Figure 8). These individuals, representing a minority within each cluster (19 individuals out of 703), seem to travel for much longer distances and probably correspond to the so-called “long travellers”, already described by Gnone et al. [37] and Carnabuci et al. [38]. All the outliers identified were excluded from subsequent analyses.
For each cluster, the single MCPs were overlapped to obtain the complex multi-polygons (MPs), which represent an approximation of the area of occupancy of the different geographical units (Figure 9). The MPs tend to occupy all of the continental platform (CP), with the clear exception of cluster E, whose MP takes only a small portion of the available CP. This is consistent with the limited (nearshore) sampling effort carried out in the corresponding study area (see also the Discussion section).

Figure 9. The multi-polygons resulting from the overlap of the single MCPs of the individuals belonging to the same cluster (see Figure 7). The MCPs were cropped following the 200 m isobaths marking the border of the continental shelf, except for cluster A (purple colour), residing in the Alboran Sea (see the main text for further details).
The difference in the average size of the MCPs across the clusters (Table 4) was found statistically significant (p-value < 0.01) based on the Kruskal–Wallis test implemented. In the pairwise comparison of the clusters, the medians were found significantly different (p-value < 0.01) in all possible combinations except for clusters A–C, D–E, D–F, and E–F (p-value > 0.05).

Table 4. The average area (km$^2$) of the individual MCP, the area (km$^2$) of the complex MP and the average width of the continental platform (CP) in the area of occupancy of the related cluster (see also Figure 9). We could not calculate the average CP width for the occupancy area of cluster F, as the Gulf of Ambracia is fully included in the continental platform.

| Cluster | Geographical Area              | N * | Average MCP (km$^2$) | Area of the MP (km$^2$) | Average Width of the CP (km) |
|---------|--------------------------------|-----|----------------------|-------------------------|-----------------------------|
| A       | Alborán Sea                    | 89  | 475.0 ± 38.4 (SE)    | 5106                    | ≈7.6                        |
| B       | Gulf of Lion–French Riviera    | 126 | 3921.2 ± 270.0 (SE)  | 18,314                  | ≈17.3                       |
| C       | Liguria–Tuscany                | 232 | 948.3 ± 50.3 (SE)    | 9384                    | ≈13.5                       |
| D       | Corsica W                      | 66  | 81.8 ± 7.8 (SE)      | 774                     | ≈3.9                        |
| E       | Corsica S–Sardinia NE          | 33  | 65.5 ± 10.2 (SE)     | 729                     | ≈18.5                       |
| F       | Gulf of Ambracia               | 113 | 98.0 ± 4.1 (SE)      | 305                     | -                           |

* Number of individual MCPs (excluding the outliers).

The average MCP and the complex MP of each cluster are positively correlated (p-value < 0.01) and seem to grow proportionally to the average width of the continental platform in the corresponding area of occupancy (Table 4): as the continental platform gets wider also the average MCP increases, following an exponential curve (Figure 10).

**Figure 10.** The average MCP and the complex MP of each cluster are positively correlated (p-value < 0.01) and seem to grow proportionally to the average width of the continental platform in the...
corresponding area of occupancy, following an exponential curve. Cluster E (whose value was not included in the regression analysis) seems to escape this general pattern, since its average MCP is quite small compared to the available platform. This is due to the limited (inshore) effort carried out in the corresponding study area (see the text for discussion on this point).

As already mentioned, cluster E seems to escape this general pattern, since both its average MCP and MP are quite small considering the available platform, but this is consistent with the nearshore sampling effort carried out in the area. In fact, most of the sightings of cluster E (565 over a total of 701, around 80%) were recorded inside the bay named “Golfo Aranci”, where dolphins are used to feed around an aquaculture facility, located a few hundred meters from the coast. The rest of the data (136 sightings, around 20%) come from the waters immediately adjacent to the gulf and from the waters between the islands of the Maddalena archipelago and the Strait of Bonifacio. Virtually no (photo-ID) data were collected in open waters, as it is clear from Figures 6, 7 and 9. We will further discuss this point in the Discussion section.

The average size of the individual MCPs and the size of the MP for each cluster are listed in Table 5. The ratio between these two values was calculated as the “overlap index”. This tends to be quite homogeneous across clusters, with the exception of cluster B (which has an overlap index about double that of clusters A, C, D, and E) and F (which has an overlap index more than triple that of clusters A, C, D, and E).

Table 5. The average area (km$^2$) of the individual MCPs and the area (km$^2$) of the multi-polygons (MPs) of each cluster. In the last column, the ratio between these two values was used to calculate the “overlap index”.

| Cluster | Geographical Area | N * | Average MCP (km$^2$) | Cluster MP (km$^2$) | Overlap Index (Average MCP/Cluster MP) |
|---------|-------------------|-----|----------------------|---------------------|----------------------------------------|
| A       | Alborán Sea       | 89  | 475.0 ± 38.4 (SE)    | 5106                | 0.09                                   |
| B       | Gulf of Lion–French Riviera | 126 | 3921.2 ± 270.0 (SE)  | 18,314              | 0.21                                   |
| C       | Liguria–Tuscany   | 232 | 948.3 ± 50.3 (SE)    | 9384                | 0.10                                   |
| D       | Corsica W         | 66  | 81.8 ± 7.8 (SE)      | 774                 | 0.11                                   |
| E       | Corsica S–Sardinia NE | 33  | 65.5 ± 10.2 (SE)     | 729                 | 0.09                                   |
| F       | Gulf of Ambracia  | 113 | 98.0 ± 4.1 (SE)      | 305                 | 0.32                                   |

* Number of individual MCPs (excluding the outliers).

3.6. Group Size

The average group size (the total number of individuals per sighting) of the clusters (Table 6) was found to be significantly different ($p$-value < 0.01) according to the Kruskal–Wallis analysis implemented. In the pairwise comparisons, the medians were found to be significantly different ($p$-value < 0.01) in all possible combinations, except for clusters D–E ($p$-value > 0.05).

Since cluster C occupies an area characterized by a large “body” on the east side, where the continental platform is quite wide, and by a long “tail” extending to the west, where the continental platform is rather narrow (see Figure 9), we also compared the average group size between these two areas. In the large portion (from La Spezia to the east end) the average group size is 15.52 (15.52 ± 0.68 SE, $n = 479$), while in the “tail” (from La Spezia to the west end) the average groups size is 13.05 (13.05 ± 0.87 SE, $n = 136$). The difference was found to be statistically significant ($p$-value < 0.05). Furthermore, the group size in the narrow portion ranges from 1 to 46 individuals, while in the large portion the group size
ranges from 1 to 100 individuals (all the large aggregations > 46 individuals were observed in the wide platform portion).

**Table 6.** The average group size (total number of individuals per sighting) in the different clusters/geographical units. In the third column the sample size (number of sightings) on which the average was calculated (n sight.).

| Cluster | Geographical Area       | n Sight. | Average Group Size |
|---------|-------------------------|----------|--------------------|
| A       | Alborán Sea             | 77       | 36.2 ± 3.64 (SE)   |
| B       | Gulf of Lion–French Riviera | 127       | 18.9 ± 1.49 (SE)   |
| C       | Liguria–Tuscany         | 615      | 14.9 ± 0.57 (SE)   |
| D       | Corsica W               | 103      | 7.7 ± 0.46 (SE)    |
| E       | Corsica S–Sardinia NE   | 696      | 4.9 ± 0.12 (SE)    |
| F       | Gulf of Ambracia        | 205      | NA *               |

* The data from the cluster F (Gulf of Ambracia) were not available for this analysis.

The “individual group sizes” (IGS, the average size of the groups in which the same individual was sighted), displayed against the individual MCPs in a scatterplot, show an interesting pattern of distribution (Figure 11), with the points well distributed along a logarithmic curve. Through the colour of the points, the data show that dolphins living in wider areas have a wider range of both the MCP and the IGS, with the clear exception of the dolphins belonging to cluster A (in the Alboran Sea), which, although having small and narrow-ranged MCP, demonstrate a high and very wide-ranging IGS. We then tested the logarithmic relationship between MCP and IGS, excluding from the analysis all the dolphins belonging to cluster A (Figure 12). The relationship was found to be statistically significant according to the ANOVA Regression test (F = 279; p-value < 0.01).

![Figure 11. The individual MCPs are displayed in the scatterplot against the individual group size (IGS). The colour of the points indicates the cluster to which they belong (the data from the cluster F—Gulf of Ambracia—were not available for this analysis).](image-url)
The correlation between the individual group size (IGS) and the individual MCP is statistically significant according to an ANOVA test ($F = 279; p$-value < 0.01). Cluster A was excluded from the analysis (see above).

We should notice that the individuals belonging to cluster E (in the geographical area Corsica S–Sardinia NE) seem to behave like dolphins from cluster D (in Corsica W), despite the inhabit an area where the continental platform is much wider. As already mentioned, this is consistent with the peculiar sampling effort carried out in the area (see also the Discussion section).

Additionally, the (logarithmic) relationship between the average group size and the average MCP across the clusters was found to be statistically significant ($p$-value < 0.01), even if only four points were displayed in the graph (see Figure 13).

![Individual MCP vs. Individual Group Size (IGS)](image1.png)

**Figure 12.** The correlation between the individual group size (IGS) and the individual MCP is statistically significant according to an ANOVA test ($F = 279; p$-value < 0.01). Cluster A was excluded from the analysis (see above).

![Average MCP vs. average group size](image2.png)

**Figure 13.** The correlation between the average MCP and the average group size across the clusters resulted statistically significant ($p$-value < 0.01). Cluster A was excluded by the analysis as in Figure 12.
3.7. Abundance Estimates and Density

The abundance estimates (only marked individuals) of the clusters/geographical units, assessed with the mark–recapture technique, are described in Table 7. In the following Table 8, the abundance estimates of the geographical units are displayed together with the size of the cluster multi-polygons (MPs), considered as an approximation of the occupancy area of the same geographical units, and the density is calculated as the ratio between these two values. As a general pattern, the size of the geographical units seems to increase with the size of the MP; however, the density shows quite different values and has its maximum value in the Gulf of Ambracia (Table 8).

Table 7. The abundance (best annual estimate, see Section 2.3.6.) of the clusters/geographical units, estimated with the mark–recapture technique.

| Cluster | Geographical Area      | Abundance (Best Estimate) of the Geographical Units |
|---------|------------------------|----------------------------------------------------|
|         | Year | n Captures | n Ind. Captured | M(t) Chao | SE | 95% CI− | 95% CI+ |
| A       | 2006 | 14         | 213             | 372       | 36.55 | 315 | 460   |
| B       | 2014 | 54         | 489             | 883       | 63.07 | 778 | 1027  |
| C       | 2011 | 100        | 296             | 503       | 45.91 | 432 | 614   |
| D       | 2007 | 57         | 98              | 107       | 5.34  | 101 | 124   |
| E       | 2009 | 47         | 37              | 114       | 55.10 | 59  | 308   |
| F       | 2012 | 41         | 103             | 115       | 6.83  | 107 | 136   |

Table 8. The abundance estimates of each geographical unit are compared with the size of the relative MPs. The density is calculated as the ratio between these two values, being the MPs considered as an approximation of the area of occupancy of the same units.

| Cluster | Geographical Area      | Abundance Est. of the Geographical Unit | Cluster MP (km²) | Density (Abundance est./Cluster MP) |
|---------|------------------------|----------------------------------------|-----------------|-----------------------------------|
| A       | Alborán Sea            | 372                                    | 5106            | 0.073                             |
| B       | Gulf of Lion–French Riviera | 883                               | 18,315          | 0.048                             |
| C       | Liguria–Tuscany        | 503                                    | 9382            | 0.054                             |
| D       | Corsica W              | 107                                    | 796             | 0.134                             |
| E       | Corsica S–Sardinia NE  | 114                                    | 729             | 0.156                             |
| F       | Gulf of Ambracia       | 115                                    | 305             | 0.377                             |

4. Discussion and Conclusions

The analyses performed in this research confirm previous findings on the distribution of the bottlenose dolphin in the context of the Mediterranean Sea: the dolphins are clustered over the continental shelf in discrete geographical units, in agreement with Gnone et al. [37] and Carnabuci et al. [38].

The physiographic characteristics of the area of residence, and specifically the size and shape of the continental shelf, seem to affect the home range of the dolphins and, consequently, the size and structure of the geographical units.

Home range and group size

The dolphins living in the Gulf of Lion, an area characterised by a wide continental shelf, are observed to have a much wider individual home range (calculated as the average MCP) compared to the units living in areas where the continental platform is very narrow,
like the west coast of Corsica (see Table 4, Figure 9). This phenomenon can in part be considered a simple “geometric” consequence of the shape and extension of the continental shelf, as it might be difficult (or impossible) for the dolphins to get a wide home range in a geographical context where the habitat boundaries are very narrow (such as the western coast of Corsica or the Gulf of Ambracia).

A direct consequence of this physical constraint seems to be the average size of the groups, as the dolphins living in those areas characterised by a narrow platform, such as the west coast of Corsica, not only show a smaller MCP, but they are also sighted in smaller groups (see Table 6, Figures 11–13). This seems quite reasonable, since a bigger group needs to exploit a bigger territory to survive [82] and home range size and group size tend to be correlated [83,84].

The average group size is small (4.9 ± 0.12 SE) also in cluster E, which inhabits a stretch of coast (Corsica S–Sardinia NE) where the continental shelf is quite large, if compared, for example, with the west of Corsica (see Table 4). However, this is not surprising considering the local sampling context. As already mentioned, the dolphins were mainly sighted (80% of all sightings) while feeding around the aquaculture fish cages of “Golfo Aranci”, a peculiar context in which the dolphins are known to form small groups. Diaz Lopez [85], who studied the bottlenose dolphins of “Golfo Aranci” during their foraging activity around the fish cages, reported an average group size of 3.5 (3.51 ± 0.1 SE). The few other sightings were recorded in the inshore waters between the islands of the Maddalena archipelago and the Strait of Bonifacio. Virtually no effort was carried out in the open waters of the continental platform, so the data available for this unit are representative of both a limited geographical area and a peculiar behavioural context (the opportunistic feeding around the aquaculture fish cages). These results support the idea that bottlenose dolphins could “fuse” into smaller groups when they enter inshore areas and/or engage in specific feeding activities, according to the “fission-fusion” mechanism described by Connor and co-authors [86].

A correlation between the group size and the “openness” of the habitat has already been reported in cetaceans and is usually considered a social strategy against predators, assuming that a larger group can better face the predation risk [87]. Our results suggest that other variables should also be taken into consideration.

In the Alboran Sea, the bottlenose dolphins form larger groups (composed by 36.2 individuals on average), but they have relatively small MCPs, according to the analysis implemented with the available data. The average MCP could have been underestimated if the research effort had been invested in an area too small to cover the movements of the dolphins in their complete extension. However, in the geographical unit of the Alboran Sea, the overlap index (see Table 5) is relatively low (0.9), meaning that the total area sampled is much wider than the average individual MCP, a result that should reassure on the reliability of the results obtained.

In this regard, it should be noted that the bottlenose dolphins living in the Alboran Sea present quite different characteristics compared to the dolphins of the other geographical units analysed. Their distribution is not confined within the continental shelf and, according to Cañadas et al. [32], these dolphins should be considered more closely related to the Atlantic offshore ecotype, which would also account for the larger size of the groups [24,26,88]. The results are not conclusive, since the distribution of the dolphins in this area could also be affected by the peculiar bathymetric profile of the Alboran Sea, characterised by a gentle slope and wide offshore shallow water shoals (such as the Alboran Ridge), which could attract the bottlenose dolphins in the open sea. However, if it were proven that the dolphins inhabiting the Alboran Sea belong to a different (offshore) ecotype, we could assume that these dolphins are able to exploit food resources in deeper bathymetric habitats, where larger groups may be more successful in their feeding activity [87,89].

It should be mentioned that the Alboran Sea presents peculiar oceanographic characteristics if compared to the inside Mediterranean. The primary production is usually much higher, thanks to the inflow of Atlantic waters through the Strait of Gibraltar, which
generates permanent anticyclonic gyres in the westernmost portion of the basin [90]. As so, some authors believe the Alboran Sea should be considered more properly as a portion of the Atlantic Ocean (at least from an oceanographic point of view), while the Mediterranean border should be moved east to the so-called Almeria-Oran front [90,91]. Additionally, as regards the cetacean fauna, the Alboran Sea shows quite a unique picture compared to the rest of the Mediterranean basin; the diversity of species is higher in this area, and the short-beaked common dolphin (Delphinus delphis), which is usually quite rare inside the Mediterranean, is the most sighted species in the shallow waters <400 m [32,33]. The presence of the common dolphin could possibly affect the behavioural ecology of the bottlenose dolphin in this sea area, shifting its preferential habitat offshore.

**Size and density**

In relation to the size of the geographical units identified, this seems to be correlated to the size of the area of occupancy (the complex multi-polygon of the cluster as a whole) which, in turn, is correlated to the size of the continental platform (Figure 10, Table 8). The bottlenose dolphin unit of the Gulf of Lion, for example, inhabiting a wide territory with no geographical or habitat breakages, was estimated to be larger than the others (see Table 7). However, when considering the density of the dolphins in their respective areas of occupancy, we found significant differences. In this regard, it should be noted that the density was estimated considering the cluster multi-polygon as an approximation of the area of residence of the related geographical unit. If the effort implemented over space was not sufficient to cover the movements of the dolphins in their complete extension, the density could be overestimated, so we should use some caution when comparing different geographical units. However, we got the maximum density (0.372, see Table 8) in the geographical units residing in the Gulf of Ambracia. Due to the peculiar physiography of this semi-closed gulf and the resident behaviour of the dolphins, this result should be considered quite reliable. In fact, the bottlenose dolphins living in the Gulf of Ambracia are considered a subpopulation, spending their entire life in the waters of the gulf [54].

It seems as though the different aggregation in bigger or smaller groups, adopted by the dolphins in the different geographical contexts, could be functional to exploit the area of residence (with its habitat constraint) to get the maximum density according to the local resources. Given this general pattern, the productivity and food resources available in the different areas of residence can obviously contribute to the density of dolphins in the same areas, as already described in both marine and land mammals [92,93].

While the size and shape of the continental shelf are stable physiographic features that can be easily measured and used as predictive parameters, the productivity and exploitability of the area of residence might change over time, according to local ecological variables, including anthropic pressures (such as the fishing effort and levy), but also according to the capability of the resident bottlenose dolphins to exploit the resources.

Behavioural plasticity is a key feature of the bottlenose dolphin and is probably transmitted from one generation to the next one as a local cultural tradition, allowing the dolphins to improve their ability to exploit the residency area and to adapt to changes. As part of this plastic behaviour, bottlenose dolphins can learn to obtain food from trawlers, gillnets and aquaculture fish cages, and these opportunistic strategies can become an integral part of their feeding habits [30,54,77,94–106].

Bottlenose dolphins can take advantage of the human activities either directly, for example, taking the fish entangled in gillnets or farmed fish [107–109], but also indirectly in the form of modified habitats that could be favourable for feeding (e.g., increase input of nutrients from aquaculture activities [99,110]). We know that human activities can have negative consequences on dolphin conservation (overfishing, bycatch, habitat degradation, noise pollution, etc.), but it can be difficult to measure the balance between negative and “positive” effects from the point of view of the bottlenose dolphins. Furthermore, the ability of bottlenose dolphins to exploit resources can change over space and time, depending on the human activities carried out locally and the behavioural adaptation of the resident
dolphins, so it may be challenging to compare the productivity of the different areas in terms of food resources actually available to dolphins.

Conclusive remarks

The results obtained in this collective research effort highlight the importance of data sharing to better understand the ecology of cetaceans in different contexts, to grasp local peculiarities, but also to understand general patterns at a higher level.

The Mediterranean bottlenose dolphin population has been recently reassessed as Least Concern by the International Union for Conservation of Nature [111], with the exception of the subpopulation inhabiting the Gulf of Ambracia, Greece, which has been assessed as Critically Endangered [112].

The Mediterranean bottlenose dolphin population is also listed in the Appendix II of the Convention on the Conservation of Migratory Species of Wild Animals (CMS), in the Appendix II of the Washington Convention on International Trade in Endangered Species (CITES), in the Appendix II of the Convention on the Conservation of European Wildlife and Natural Habitats (Bern Convention), in Appendix II of the Protocol to the Barcelona Convention on Specially Protected Areas of Mediterranean Importance (SPAMI). Finally, the bottlenose dolphin is included in the Annexes II and IV of the EU Habitats Directive (Council Directive 92/43/EEC) as a species of “community interest whose conservation requires both designation of special areas and strict protection” [113].

Our findings can help to better characterise the different geographical and management units that constitute the Mediterranean bottlenose dolphin metapopulation in relation to the physiographic characteristics of their area of residence. This can contribute to develop proper management and conservation measures for the bottlenose dolphin in the context of the Mediterranean Sea and beyond.

At the same time, further insights are needed to complete the picture. For example, we should consider that most of the data analysed were collected in daylight time and in the favourable conditions (i.e., the calm sea and long daylight time) given by the “good season” (late spring, summertime, early autumn). We cannot exclude that the behavioural patterns of the bottlenose dolphins, in relation to the continental platform, could change during night time or in winter. Further studies, possibly including other techniques such as passive acoustic, could help to monitor the behaviour of the dolphins in these other contexts.

Furthermore, the role of so-called “long travellers” needs to be better clarified. These dolphins, despite being a clear minority, could represent a means of continuity between the different geographical units, and their role in this regard could be more important than is understood.

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