Population Dynamics Reveal a Core Community of the Common Bottlenose Dolphin (*Tursiops truncatus*) in Open Waters of the South-Western Gulf of Mexico

Jaime Bolaños-Jiménez1,2, Eduardo Morteo2,3*, Christian A. Delfín-Alfonso2,3, Pedro F. Fruet4,5,6, Eduardo R. Secchi6 and Javier Bello-Pineda1

1 Instituto de Ciencias Marinas y Pesquerías (ICIMAP), Universidad Veracruzana, Veracruz, Mexico, 2 Laboratorio de Mamíferos Marinos (LabMMar-IIB-ICIMAP), Universidad Veracruzana, Veracruz, Mexico, 3 Instituto de Investigaciones Biológicas, Universidad Veracruzana, Xalapa, Mexico, 4 Museu Oceanográfico Prof. Eliézer de C. Rios, Universidade Federal do Rio Grande/FURG, Rio Grande, Brazil, 5 Kaos, Rio Grande, Brazil, 6 Laboratório de Ecologia e Conservação da Megafauna Marinha (ECOMEGA), Instituto de Oceanografia, Universidade Federal do Rio Grande/FURG, Rio Grande, Brazil

The presence of transient and temporary individuals in capture-mark-recapture studies may violate the assumption on equal catchability, and thus yield biased estimates. We investigated the effects of residency patterns on population parameters of bottlenose dolphins inhabiting the coastal waters off the Alvarado Lagoon System (ALS), Veracruz, Mexico. We hypothesized that this population is open but there exists a “core community” that behaves as a closed population. Between 2006 and 2010, we conducted 75 photo-identification surveys and recorded 263 dolphin group encounters, in which 231 dolphins were identified. Individuals present during only one season, classified as transients (*n* = 85), were excluded from the study, and a standardized residency index (*IH*4) was computed for each dolphin that remained in the sample (*n* = 146). We used the K-means clustering method to split the sample into groups based on individual (seasonal, annual) *IH*4 values. These clusters were named as regular residents (RR, *n* = 55), occasional residents (OR, *n* = 45), and occasional visitors (OV, *n* = 46). The cumulative frequency of newly identified individuals displayed an asymptotic trend for the whole sample and all clusters, indicating that most of the individuals present in the study area during the study period were identified. The assumption of demographic closure was tested to define the core community, and was rejected for the whole sample and the OV cluster (*p* < 0.001 in both cases), indicating that the population is open. The closure assumption was not rejected for RR and OR clusters (*χ*2 = 6.88, DF = 13, *p* = 0.91, and *χ*2 = 17.8, DF = 16, *p* = 0.33, respectively), indicating that these clusters were demographically closed over the 5-year period. Thus, we defined this aggregation of individuals as the “core community”. The closed population model *M*1 indicated that the total abundance of this core community was 123 individuals (95% CI: 114–133). Our results provide quantitative evidence of the
INTRODUCTION

In most wildlife populations, sex and age have profound effects on the chances of an individual dying, or producing offspring (Neal, 2004), thus these factors are routinely considered when studying population structure and dynamics (Ojasti and Dallmeier, 2000). Other factors with the potential to influence population structure are site fidelity (SF) and philopatry (Bose et al., 2017). SF has been defined as the “tendency to return to a previously occupied location” (Greenwood, 1980; Switzer, 1993), whereas philopatry is the tendency to stay in a familiar environment, mainly related to the natal habitat (Begon et al., 1995). Several environmental factors have been suggested to influence species’ site fidelity and/or philopatry, such as habitat stability, predictability of reproductive failure, variability in territory quality within a habitat, and population pressure; individual characteristics may also include an individual’s previous reproductive success, age, and knowledge of other sites (Switzer, 1993).

In cetacean studies, SF and residency patterns are evaluated using information on the number of recaptures, the duration of stay, and/or the average recurrence of individuals (see Ballance, 1990 and Morteo et al., 2012b). In this regard, Ballance (1990), stated that “a high number of resightings, a long period time between the first sighting and the last resighting, and a short time interval between adjacent sightings describes an animal with a high degree of residence.” Thus, individual residency patterns can be considered as a manifestation of SF and philopatry.

For cetacean social species such as delphinids, other factors that are thought to shape population and social structure are kinship and reproductive condition (Bigg, 1982; Wells et al., 1987). Sex and age also influence the way that delphinids use their habitats. In general, females try to ensure the survival of their offspring, thus their distribution is often closely related to the quality and/or accessibility of food resources and habitats that have a lower risk of predation; conversely, the distribution of males is often related more to female distribution than to the availability of food (Gowans et al., 2007). Delphinids routinely travel throughout the day and can be exposed to large areas, their entire home range, or even multiple habitats on a daily basis (Gowans et al., 2007). Consequently, individual home ranges - that may depend on the age and sex of the individual- result in structured societies that are usually stable over time (Wells et al., 1980, 1987), in which females tend to be more resident to specific areas than males (Gowans et al., 2007).

Since Ballance’s (1990) work, several methods have been used to evaluate cetacean residence and this lack of standardization hinders proper comparison among SF studies (Tschopp et al., 2018, see below). Tschopp et al. (2018) found that, in general, the methods to quantify site fidelity include three approaches: (1) proportions (a ratio between the number of sightings or resightings, and a measure of effort), (2) categories (e.g., high, moderate, low), and (3) models (maximum likelihood methods). Recently, Tschopp et al. (2018) developed and compared the performance of a series of standardized indexes -based on Ballance’s parameters- to be able to quantify the site fidelity degree and ensure accurate comparability across related investigations. Regarding delphinids, several authors (e.g., Gowans et al., 2007; Wells and Scott, 2018) have predicted that in offshore environments, where food availability is patchy and unpredictable and predation risk might be high, dolphins range more widely and form larger groups to forage on sparsely distributed prey schools and to reduce predation. In contrast, in complex inshore environments with predictable resource availability and potentially lower predation, dolphins are predicted to remain resident in relatively small areas and to form smaller groups to avoid feeding competition. This seems to be the case for the bottlenose dolphin (Tursiops truncatus) populations inhabiting semi-enclosed bays or sounds, or inshore habitats in the Gulf of Mexico (GoM). This species has been widely studied in the northern GoM on the basis of photo-ID methods and capture-mark-recapture (CMR) models (see review in Vollmer and Rosel, 2013). In the northern GoM, where long-term, year-round residents are best documented, there is also evidence for seasonal changes in abundance; most often these are thought to result from movements of seasonal, short-term residents and/or transients (very short-term visitors) to the area. It is generally thought that these shorter-term residents and visitors come from the adjacent coastal population, although comprehensive and directed studies to address this question have yet to be performed (Vollmer and Rosel, 2013).

The bottlenose dolphin has been widely documented in some localities in the southern GoM (Ortega Ortiz, 2002; Ortega Ortiz et al., 2004; Ramirez-León et al., 2020), including waters off the Tamiahua Lagoon System (Galindo et al., 2009; Martínez-Serrano et al., 2011; Valdés-Arellanes et al., 2011); Nautla (Morteo et al., 2019); and the Alvarado Lagoon System (ALS) in Veracruz state (Morteo, 2011; Morteo et al., 2012b, 2014, 2017, 2019; Morales-Rincón et al., 2019), and some coastal areas in the states of Tabasco, Campeche and Yucatán (Delgado-Estrella, 2015).

Off the ALS, Morteo et al. (2014) found that this population is sexually segregated, where females were more resident, had higher site fidelity, and had weaker associations with a higher number of partners than males. On the other side, males were assumed to be primarily responsible for gene flow among adjacent locations (Morteo et al., 2014).
Capture-mark-recapture (CMR) studies have been used as a general sampling and analysis method to assess population status and trends in many biological populations (White and Burnham, 1999). The photographic documentation of long-lasting natural marks on the dorsal fin or flukes (i.e., photo-identification, photo-ID) has been used since the early 1970’s to study aspects such as group structure, site fidelity, movement patterns, and abundance using CMR models in cetaceans (Würsig and Jefferson, 1990). Models usually require a set of restrictive assumptions about the properties of the population under study, such as individual homogeneity in capture and survival probabilities, among others (Krebs, 1972). Failure to comply with these assumptions implies that the model does not adequately fit the data, and thus may introduce severe bias in parameter estimates (Krebs, 1972).

In structured delphinid populations, heterogeneity in capture or survival probabilities is often produced by age, sex, or size of the individuals (Williams et al., 1993), as well as inconspicuous marks, low photographic quality, and social bonds (Morteo et al., 2012a).

In recent studies, researchers started to include site fidelity and residency pattern as factors to account for heterogeneity in the data when studying demographics or population ecology of several cetacean species, including the southern Australian bottlenose dolphin (Tursiops cf. australis, Zanardo et al., 2016; Passadore et al., 2017), the Australian humpback dolphin (Sousa sahulensis, Hunt et al., 2017), the Risso’s dolphin (Grampus griseus, Carlucci et al., 2020) and the fin whale (Balaenoptera physalus, Schleimer et al., 2019). These researchers calculated sighting rates and site fidelity indexes for each individual and then used agglomerative clustering methods (Legendre and Legendre, 1998) to identify clusters of individuals with similar degrees of site fidelity, before running CMR analysis. In particular, Haughey et al. (2020) used for the first time Tschopp et al. (2018) standardized indexes to stratify the sample and estimate population parameters of Indo-Pacific bottlenose dolphin (T. aduncus) off western Australia by residency pattern.

In this paper, we used the recently developed standardized site fidelity index (SSFI) IH4 (Tschopp et al., 2018) to study potential differences in population parameters between resident and non-resident common bottlenose dolphins (Tursiops truncatus) that use the open, coastal waters off Alvarado Lagoon System, southwestern Gulf of Mexico. On the basis of our knowledge of the study area, we hypothesized that bottlenose dolphins using the marine coastal waters adjacent to the ALS are part of an open population within which exists a core community (sensu Wells et al., 1987), that behaves as a closed population, in the sense that it is composed of all the groups and individuals “… that share large portions of their ranges and interact with each other to a much greater extent than with members of similar units in adjacent waters, but genetics exchange occurs between communities.”

MATERIALS AND METHODS

For this study, we used the photo-ID catalog and database from the Marine Mammal Lab at Universidad Veracruzana (LabMMar-IIB-ICIMAP). This database includes the sighting histories of 231 bottlenose dolphins from 75 survey trips conducted between May 2006 and August 2010 (Morteo, 2011; Morteo et al., 2012b, 2014, 2017). Previously, Morteo et al. (2014, 2017) used a partial subset of these databases, covering the period May 2006–April 2008, to explore the abundance and social ecology of this dolphin population under open population modeling by considering the population as a homogeneous unit, and without considering temporal variations or population structure in the parameters. Detailed definitions of methodological terms are presented in Supplementary Material 1.

Study Area

The coastal waters of Alvarado, Veracruz, are shallow (less than 20 m), with an average temperature of 27°C (Morteo, 2011), and are strongly influenced by the discharge of the Alvarado Lagoon System (ALS, de la Lanza Espino and Lozano Montes, 1999; Cruz-Escalona et al., 2007). The ALS is a coastal wetland located in the center-south of the State of Veracruz, southwestern Gulf of Mexico, formed by the confluence of the Acula, Blanco, Limón, and -mainly- the Papaloapan rivers (Figure 1). The ALS has an elongated shape, parallel to the coastline, with an approximate length of 26 km, with a maximum width of 5 km and an average depth of 2.5 m, for a total area of about 80 km² (de la Lanza Espino and Lozano Montes, 1999). The weather is tropical, with three marked climatic seasons; following Morteo (2011) and Morteo et al. (2014), we defined these seasons as: Dry, with a significant reduction in average precipitation, from March to June; Rainy, in which runoff causes high organic matter and nutrient input into coastal waters, from July to October; and, finally, the “Nortes” or “Northern Winds,” with strong winds associated to cold fronts which may last several days (de la Lanza Espino and Lozano Montes, 1999; Cruz-Escalona et al., 2007).

Survey Protocol

The survey protocol represents an extension of the procedures described in Morteo (2011), and Morteo et al. (2014, 2017). All sampling surveys were conducted following a zigzag pattern that started at the mouth of the ALS and extended 4 km offshore, and 9 km on each side of the estuary mouth (Morteo, 2011; Morteo et al., 2014, 2017, see Figure 1). Typically, all surveys covered the whole study area, and no surveys were conducted inside the lagoon system, based on previous data showing virtually no presence of dolphins (Morteo, pers. obs.).

Dolphin Photo-ID

Standard photo-ID techniques were used to individually identify and catalog adult dolphins based on long-lasting marks in their dorsal fins (Würsig and Jefferson, 1990; Morteo et al., 2014, 2017; Urian et al., 2015). All photographs were graded for quality (PQ) and dolphin distinctiveness (D). Following Urian et al. (2013, 2015), only excellent (PQ1) and good quality (PQ2) pictures of very (D1) and average (D2) distinctive dolphins were used for analysis, to minimize misidentification. We then plotted discovery curves
for new identified individuals across time (Fisher et al., 1943; Colwell et al., 2004).

**Mark Rate**

A mark rate ($\theta$), representing the proportion of marked animals in the groups, was calculated for each sighting to adjust abundance estimates by accounting for the unmarked fraction of the population, and thus produce estimates of seasonal abundances for the full population (Wilson et al., 1999; Wickman et al., 2020). Following Morteo et al. (2014, 2017), we used a combination of field and analytical approaches to accurately estimate group size and $\theta$ for each encounter. In the field, minimum, maximum, and best group size estimates were recorded, and the best estimate was used to calculate a mark rate for each sighting. By using the full dataset [$N = 263$ groups], we found that the average difference between the minimum and maximum group size estimates was 0.92 dolphins ($\pm 3.70$ SD) and this difference increased with larger groups. As the average group size for the full study period was fairly small ($9.0 \pm 11.2$ SD) and the standard deviation of the difference between extreme field estimates was less than one individual, we considered that our group size estimates are highly accurate, and thus representative of the total number of animals in the group. Additionally, we used the empirical criteria developed independently by Würsig (1978) and Ballance (1990) to double-check dolphin counts on the field; this approach states that the probability of having photographed all the dolphins in a group is higher than 95% if all marked individuals in the sighting were correctly photographed at least four times (Ballance, 1990; Würsig and Jefferson, 1990).

**Residency Pattern Assessment**

Morteo et al. (2012b) reviewed the parameters proposed by Ballance (1990), suggested modifications, and established “Occurrence,” “Permanence” and “Periodicity,” as their names. The occurrence was redefined as the number of times the animal was recaptured (that is, eliminating the first sighting from the calculation); the permanence was defined as “the time over which an individual was recorded, determined by the difference between its first and last sighting,” whereas the periodicity was redefined as “the recurrence of the individual, determined by the inverse of the average time (in days) between consecutive recaptures.” Tschopp et al. (2018) used these definitions to evaluate the performance of their standardized indexes. For the purpose of this study, the periodicity was calculated by season and year.

We used the IH$_4$ index of Tschopp et al. (2018) to split the sample into clusters according to individual’s site fidelity. We chose this index because it consistently had the best performance in all of the scenarios, and the authors proposed it as a standardized measure of site fidelity (SSFI). The IH$_4$ is based on the harmonic mean of parameters permanence (IT) and
periodicity (It), as expressed in equation 1:

\[
IH_4 = \frac{2}{IT + \frac{n}{T}}
\]  

(1)

Where: IT = Permanence  
It = Periodicity

For each dolphin, we calculated the IH₄ per season and year, to account for their presence in the study area across the studied period under different temporal stratifications. We then constructed a matrix (IH₄-matrix) in which each line corresponded to an individual dolphin and columns included the two IH₄ measures. We used the IH₄-matrix to split the sample into groups by using the K-means clustering method (Legendre and Legendre, 1998). With the K-means method, a set of n objects (= dolphins) in a p-dimensional space (= IH₄-matrix) can be partitioned into K groups -or clusters- such that the objects within each cluster are more similar to one another than to objects in other clusters; the number of groups (K) is often determined by the user based on expert knowledge (Legendre and Legendre, 1998), but there exist some numeric criteria to select the best grouping strategy (see below). The function cascade KM of the “vegan” package (Oksanen et al., 2019) of the R environment (R Core Team, 2020) was used to determine the groupings with the K-means method. The best-fitting K was selected by means of the Calinski-Harabasz index (ICH, Calinski and Harabasz, 1974; Legendre and Legendre, 1998) available in the vegan package. According to the ICH, the grouping with the highest index value corresponds to the best grouping given the data (Calinski and Harabasz, 1974; Oksanen et al., 2019). As we were interested in population analysis by the residency pattern, we excluded the transient individuals from the sample. We defined as transients those individuals that were present in the study area only during one season, independently of the number of recaptures during that season. Through the cluster analysis, we then compared the performance of 2 and 3 clusters (K), respectively.

### Population Parameters

Estimates were made under standard CMR models in the program MARK (White and Burnham, 1999). Parameters can be established as time-variant or time-invariant, as well as variant or invariant among groups. Following White and Burnham (1999), the notation (.) was used to indicate time-variant parameters and (t) to indicate those time-variant parameters. The notation (g) was used to indicate time-invariant parameters, different for each residency group, and (g^t) to indicate parameters different for each group and time-variant. The model(s) that best fitted the data were selected by using the lowest value of the Akaike Information Criterion, corrected for small samples (AICc, Akaike, 1973; White and Burnham, 1999). Before analyses, sighting histories were collapsed by season and stratified by residency pattern. We used two steps to estimate population parameters. In the first step, we used the POPAN superpopulation approach of the Jolly-Seber model (Schwarz and Arnason, 1996; White and Burnham, 1999) to determine survival, recruitment, and seasonal abundance of the dolphins according to their residency pattern under the assumption of open population. In the second step, we used closed capture-recapture methods available in the CAPTURE routine of the MARK program to determine the seasonal abundances of the core community only. Finally, total abundances were estimated as:

\[
N_T = \frac{N_m}{\theta}
\]  

(2)

where Nₜotal = total abundance, Nₘ = abundance of marked individuals, and θ = mark rate. The variance was calculated by using the Delta method (Wilson et al., 1999) as:

\[
Var(N_T) = N_T^2 \left[ \frac{var(N_m)}{N_m^2} + \frac{1-\theta}{n\theta} \right]
\]  

(3)

and then, log-normal 95% confidence intervals (Burnham et al., 1987; Tezanos-Pinto et al., 2013) were calculated as:

\[
N_{lower} = \frac{N_T}{C} \quad \text{and} \quad N_{upper} = N_T \times C
\]

where Nₜₗₜₜ and Nₜᵤₜₑₑₑ are the lower and upper bound, respectively, of the confidence interval, and

\[
C = z_{0.025} \times \sqrt{\log_e \left[ 1 + CV(N_T)^2 \right]}
\]  

(5)

where z₀.₀₂₅ is the normal deviate (1.96), and CV is the coefficient of variation.

### Assumption Compliance and Parameter Estimations

Valid inference in CMR studies requires compliance with several assumptions (Pollock et al., 1990; White and Burnham, 1999). The U-CARE software (version 2.3.2, Choquet et al., 2009, 2005) was used to assess the goodness of fit of the model(s) to the data. We used the POPAN parameterization of the Jolly-Seber model (Schwarz and Arnason, 1996) to get estimates for each group of apparent survival (Phi, hereafter survival), catchability (p), and probability of entrance of individuals from the superpopulation (hereafter recruitment) to the study area. Survival is termed “apparent” as the algorithm cannot discriminate between mortality and permanent emigration (White and Burnham, 1999). We used the Closed Population approach (Otis et al., 1978; Chao et al., 1992; Stanley and Burnham, 1999) to get estimates of the population size for the more resident fraction of the population. The closure assumption was confirmed with the closure test by Stanley and Burnham (1999), as provided by the Close Test software (Stanley and Burnham, 1999; Stanley and Richards, 2005). We applied these tests to the whole sample (n = 231 individuals), as well as to the clusters obtained through the k-means method. In all cases, the probability of first capture was set equal to the probability of recapture (p = c), as the process of taking photographs does not involve the physical capture of individuals so no behavioral effect is expected (Tezanos-Pinto et al., 2013).
Between May 2006 and August 2010, we conducted 75 surveys, totaling 6,328.8 km surveyed, 288.1 h of search effort, and 110.4 h photographing dolphin groups (Table 1). Search effort ranged from 3.8 to 8.15 h day$^{-1}$ ($\bar{x} = 3.6 \pm 1.1$ SD) and the full study area was covered in each and all survey trips. In general, the effort was greater during the first two seasons, but we found no evidence of a difference in search effort among seasons (Kruskal-Wallis, $H = 0.91, p = 0.63$). Photo-ID effort (h day$^{-1}$) ranged from 0.0 to 3.6 (mean $1.4 \pm 0.8$ SD), and no seasonal differences were found (Kruskal-Wallis, $H = 10.55, p = 0.10$).

Two hundred and sixty-three groups were encountered totaling 2,320 dolphins sampled, of which 231 distinct individuals were identified. We processed 30,402 pictures of which 10,958 (36%) were useful for photo-ID. Group size averaged 9.0 individuals ($\pm 11.2$ SD, range: 1–100), and remained similar among seasons (Kruskal-Wallis, $H = 2.62, p = 0.27$). The number of recaptures for marked dolphins ranged from 0 to 12 occasions, where 36.8% ($n = 85$) were transients (Figure 2).

### Clustering of Individuals

The ICH (Calinski and Harabasz, 1974) indicated that $K = 3$ (ICH = 389.4) was the best grouping strategy according to the data. Cluster 1 consisted of 46 individuals with low seasonal (mean = $0.27 \pm 0.08$ SD) and annual (mean = $0.31 \pm 0.18$ SD) site fidelity indexes. These individuals were resighted between 1 and 4 seasons. Cluster 2 comprised 45 individuals with medium seasonal (mean = $0.49 \pm 0.10$ SD) and annual (mean = $0.76 \pm 0.10$ SD) site fidelity indexes; these individuals were resighted between 2 and 8 seasons. Cluster 3 consisted of 55 individuals with high seasonal (mean = $0.79 \pm 0.09$ SD) and annual (mean = $0.96 \pm 0.06$ SD) site fidelity indexes; these individuals were recaptured between 5 and 12 seasons. Following Zanardo et al. (2016) and Passadore et al. (2017) these clusters were defined as “occasional visitors” (OV), “occasional residents” (OR), and “regular residents” (RR), respectively (Figures 3, see definitions in Supplementary Material 1).

The Kruskal-Wallis test showed that there were differences in the medians of the IH$_{4}$ values among groups (IH$_{4}$-season: $p < 0.001$, IH$_{4}$-year: $p < 0.001$). The Mann-Whitney pairwise post-hoc test showed that the medians of the groups were different (IH$_{4}$-season: $p < 0.001$; IH$_{4}$-year: $p < 0.001$, in all cases), indicating differences in the time spent in the study area for individuals classified in different clusters.

### Discovery Curves

The cumulative frequency of newly identified individuals by residency pattern displayed an asymptotic trend in all cases (Figure 4), indicating that most of the individuals present in the study area for the duration of the study were identified. For RR and OR individuals, the curve reached a plateau within the first seasons of our study. Additionally, for OV, transients (TR), and

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**TABLE 1** Summary of survey and photo-identification effort of bottlenose dolphins off Alvarado Veracruz System, south-western Gulf of Mexico.

| Season  | n  | $sT$ (hs) | $pT$ (hs) | $nE$ | km  |
|---------|----|-----------|-----------|------|-----|
| Dry-2006| 4  | 22.9      | 5.5       | 19   | 368.6 |
| Rainy-2006| 13 | 56.3      | 25.1      | 48   | 1031.6 |
| NW-2007 | 5  | 20.4      | 7.4       | 25   | 357.3  |
| Dry-2007| 7  | 27.3      | 10.0      | 25   | 519.9  |
| Rainy-2007| 6  | 23.5      | 9.4       | 18   | 512.1  |
| NW-2008 | 2  | 7.1       | 1.3       | 7    | 131.6  |
| Dry-2008| 2  | 10.9      | 2.1       | 6    | 214.9  |
| Rainy-2008| 4  | 11.9      | 8.6       | 17   | 308.3  |
| NW-2009 | 5  | 20.6      | 9.6       | 21   | 658.4  |
| Dry-2009| 8  | 27.4      | 9.5       | 30   | 797.7  |
| Rainy-2009| 7  | 20.2      | 6.6       | 15   | 463.3  |
| NW-2010 | 3  | 12.3      | 2.7       | 6    | 308.4  |
| Dry-2010| 6  | 20.9      | 9.9       | 20   | 515.0  |
| Rainy-2010| 3  | 6.6       | 2.8       | 6    | 141.6  |
| Totals  | 75 | 288.1     | 110.4     | 263  | 6328.8 |

$n = \text{No. of surveys}, sT = \text{Search effort}, pT = \text{Photo-ID effort}, nE = \text{No. of dolphin group encounters}, km = \text{km surveyed}.$
FIGURE 3 | Boxplot of IH₄ values, by season and year, according to the residency pattern. OV = occasional visitors (n = 46), OR = occasional residents (n = 45), RR = regular residents (n = 55).

FIGURE 4 | Discovery curve of the cumulative frequency of newly identified bottlenose dolphins off the Alvarado Lagoon System, between 2006 and 2010, by residency pattern. FS = full sample, TR = transients, RR = regular residents, OV = occasional visitors, OR = occasional residents, NW = Northern Winds. Number of individuals belonging to each cluster are indicated.
the whole sample, the curves presented two plateaus, indicating at least one occasional pulse of incorporation of individuals to the study area (Figure 4).

**Goodness of Fit Tests and Model Selection**

The global test for the sample stratified by residency pattern (excluding transients) was non-significant ($\chi^2 = 70.1$, D.F. = 78, P-value = 0.73), indicating compliance with the model assumptions.

The most parsimonious model (AICc = 1810.9; AICc weight = 0.83) indicated that survival was time-invariant, different among groups, and that both catchability and recruitment were time-variant and group-variant [$\Phi(g)$, $p(g^{*t})$ pent$(g^{*t})$, model 1 in Table 2]. Formulations with invariant recruitment [$pent(g)$, pent(.) or equal for all groups pent(t)] failed to converge or presented unrealistic values or null standard error, thus they were not included in the analyses.

**Population Parameters**

Catchability (Figure 5) for RR (mean 0.67 ± 0.25 SD) was higher than for OR (0.35 ± 0.20 SD) and OV (mean 0.33 ± 0.27 SD) (Anova, $F = 8.64, p = 0.0008$).

**Apparent Survival and Recruitment**

According to the most parsimonious model, seasonal survival was time-invariant and resulted in 1.00 (95% CI: 1.00–1.00) for RR and OR, and 0.91 (95% CI: 0.85–0.95) for OV individuals. When scaled to represent annual rates (see Cooch and White, 2019, p. 4.27), survival was 1.00 (95% CI: 1.00–1.00) for RR and OR, and 0.75 (95% CI: 0.61–0.86) for OV. Recruitment for the duration of the study was 0.72, 0.20, and 0.02 for OV, OR, and RR clusters, respectively, such that 28%, 80%, and 98% of OV, OR, and RR individuals, respectively, were already present in the population just before the beginning of the study (Table 3).

Occasional visitors were recruited in up to five out of 14 sampling occasions, with peaks in the rainy season of 2006 (23%), and the northern winds season of 2009 (30%). Occasional residents were recruited only during the NW 2007 and Dry 2008 seasons, with the highest percentage in the dry season of 2007 (16%). Regular residents were recruited only in the Rainy season of 2007, but in a low proportion (2%).

The high survival values of the RR and OR clusters, as well as the null and low recruitment rates of RR and OR individuals, respectively, were consistent with the assumption of demographic or geographical closure for these clusters. As stated earlier, additional analyses were conducted to determine abundance for the core community, under the assumption of closed populations (see below).

**Abundance**

All abundances were adjusted by $\theta$ to include the unmarked fraction of the population. The total abundance of occasional visitors averaged 34 individuals (± 6.0 D.E., range: 18–42). The abundance of occasional residents averaged 57 (± 7.3 D.E., range: 44–76), and regular residents averaged 72 (± 7.2 D.E., range: 65–93) dolphins. A regression analysis indicated a low, positive, non-significant trend in the seasonal abundance of OV individuals ($p = 0.09, R^2 = 0.21$). The residual plot of abundances for all clusters showed a non-random pattern (see Supplementary Material 2).

**Closed Population Approach**

The closure assumption was rejected for the whole sample and the OV cluster ($p < 0.001$ in both cases), however, this was not the case for the RR and OR clusters ($\chi^2 = 6.88, DF = 13, p = 0.91$, and $\chi^2 = 17.8, DF = 16, p = 0.33$, respectively). Thus, we defined the aggregation of these clusters as the “core community” (Wells et al., 1987), and then used closed models in MARK to determine its abundance by means of the $M_{th}$ formulation (Chao et al., 1992), which includes heterogeneity (Chao et al., 1992; White and Burnham, 1999; Cooch and White, 2019). We chose the $M_{th}$ estimator (Chao et al., 1992) because the sampling protocol does not require the animals to be physically captured, and because of the heterogeneity in the capture probabilities of the OR and RR clusters (see section “Population parameters”). According to the $M_{th}$ formulation, the abundance of marked adult individuals (RR+OR) was 100 individuals (SE = 0.21). When corrected for the average mark rate (0.81 ± 0.17 SD), total abundance of the core community was 123 individuals (95% CI: 114–133). Interestingly, the closure test for the “core community” (OR+RR) was rejected ($\chi^2 = 49.5, DF = 16, p = 0.00003$), indicating a potential violation of this assumption; nevertheless, as both OR and RR clusters separately complied with the closure assumption, we believe that the significant result is an artifact of the different capture probabilities of each cluster (0.35 and 0.67, respectively). In this regard, it is well-known that heterogeneity in capture probabilities can cause both the tests of immigration and emigration to reject the null hypothesis of closure (Cooch and White, 2019, p. 14.6).
TABLE 2 | Set of candidate models arranged in ascending order by AICc for population analysis of the bottlenose dolphin off Alvarado Veracruz System, southwestern Gulf of Mexico, between 2006 and 2010.

| Model | AICc | Delta AICc | AICc weights | Model likelihood | Number of parameters | Deviance | −2log(L) |
|-------|------|------------|--------------|------------------|---------------------|----------|----------|
| 1 (\Phi(g) p(g\text{*}) p\text{ent}(g\text{*})) | 1810.9 | 0.0 | 0.83 | 1.00 | 87 | 544.3 | 1616.5 |
| 2 (\Phi(\text{ }) p(g\text{*}) p\text{ent}(g\text{*})) | 1814.7 | 3.8 | 0.13 | 0.15 | 85 | 553.1 | 1625.2 |
| 3 (\Phi(t) p(g\text{*}) p\text{ent}(g\text{*})) | 1816.9 | 6.0 | 0.04 | 0.05 | 84 | 557.8 | 1629.9 |
| 4 (\Phi(g) p(t\text{*}) p\text{ent}(g\text{*})) | 1977.7 | 166.8 | 0.00 | 0.00 | 59 | 778.4 | 1850.6 |
| 5 (\Phi(g\text{*}) p(t\text{*}) p\text{ent}(g\text{*})) | 1986.2 | 175.2 | 0.00 | 0.00 | 81 | 734.4 | 1806.6 |
| 6 (\Phi(t) p(t\text{*}) p\text{ent}(g\text{*})) | 2035.1 | 224.2 | 0.00 | 0.00 | 60 | 833.5 | 1905.7 |
| 7 (\Phi(t\text{*}) p(t\text{*}) p\text{ent}(g\text{*})) | 2049.5 | 238.6 | 0.00 | 0.00 | 57 | 854.8 | 1927.0 |
| 8 (\Phi(g\text{*}) p(g) p\text{ent}(g\text{*})) | 2055.6 | 244.7 | 0.00 | 0.00 | 83 | 798.9 | 1871.1 |
| 9 (\Phi(t) p(g) p\text{ent}(g\text{*})) | 2057.4 | 246.5 | 0.00 | 0.00 | 58 | 860.4 | 1932.6 |
| 10 (\Phi(\text{ }) p(g) p\text{ent}(g\text{*})) | 2088.3 | 277.4 | 0.00 | 0.00 | 48 | 914.2 | 1986.4 |
| 11 (\Phi(t) p(g) p\text{ent}(g\text{*})) | 2126.0 | 315.1 | 0.00 | 0.00 | 46 | 956.4 | 2028.6 |
| 12 (\Phi(g) p(t) p\text{ent}(g\text{*})) | 2174.2 | 363.2 | 0.00 | 0.00 | 13 | 1075.6 | 2147.7 |
| 13 (\Phi(g\text{*}) p(t) p\text{ent}(g\text{*})) | 2208.8 | 397.9 | 0.00 | 0.00 | 82 | 954.6 | 2026.8 |
| 14 (\Phi(t) p(t) p\text{ent}(g\text{*})) | 2240.9 | 429.9 | 0.00 | 0.00 | 56 | 1048.5 | 2120.7 |
| 15 (\Phi(t\text{*}) p(t) p\text{ent}(g\text{*})) | 2251.2 | 440.3 | 0.00 | 0.00 | 13 | 1152.6 | 2224.8 |
| 16 (\Phi(g\text{*}) p(g\text{*}) p\text{ent}(g\text{*})) | 57050.9 | 55239.9 | 0.00 | 0.00 | 116 | 55709.0 | 56781.2 |

Sample stratified by residency pattern.

DISCUSSION

Population structure of highly mobile marine organisms can be complex and difficult to study, but it is important to understand how individuals within a population partition their environment, in order to better address conservation challenges (Vollmer et al., 2021). In this regard, recent advancements in the implementation of standardized site fidelity indexes (Tschopp et al., 2018) and considerations on residency patterns when studying cetacean population dynamics (e.g., Zanardo et al., 2016; Hunt et al., 2017; Passadore et al., 2017; Schleimer et al., 2019; Carlucci et al., 2020; Haughey et al., 2020) provide improved methodological tools for dealing with heterogeneity in capture or survival probabilities inherent to the dynamics of social species with fission-fusion societies such as bottlenose dolphins (Connor et al., 2000). This is the first study to explore the population dynamics of the common bottlenose dolphin (T. truncatus) in the Gulf of Mexico based on its residency patterns, and to provide quantitative evidence of the existence of a “core community” (sensu Wells et al., 1987) in an open, coastal habitat. As this study is an extension of previous work by Morteo (2011) and Morteo et al. (2014, 2017), we included new data on previously unknown parameters, improving the temporal coverage as well as a more comprehensive analytical approach that yielded more refined estimates, and a more robust assessment of the population structure.

TABLE 3 | Recruitment from the superpopulation of the bottlenose dolphin off the Alvarado Lagoon System, southwestern Gulf of Mexico, 2006–2010.

| Season       | Occasional visitors | Occasional residents | Regular residents |
|--------------|---------------------|----------------------|------------------|
| Rainy 2006  | 0.23 (0.12–0.41)    | 0.16 (0.02–0.57)    |                  |
| NW 2007     | 0.14 (0.03–0.44)    | 0.02 (0.00–0.12)    |                  |
| Dry 2008    | 0.03 (0.00–0.09)    | 0.03 (0.00–0.62)    |                  |
| Dry 2009    | 0.02 (0.00–0.09)    | 0.03 (0.00–0.62)    |                  |
| Dry 2010    | 0.30 (0.14–0.52)    | 0.02                 |                  |

95% CI’s are indicated between parentheses. Occasions with null recruitment are not indicated. NW = northern winds.
Due to the longevity of bottlenose dolphins, we acknowledge that short-term studies such as this one may not encapsulate the population processes related to factors such as mortality or emigration (Haughey et al., 2020). Despite the high number of transient individuals in the sample (Figure 2), the asymptotic trend in the discovery curves (Figure 4) indicated that most of the marked individuals in the population had been photographed by the end of the study period. For the full sample, the existence of two plateaus and decrease in the discovery rate of new individuals after the rainy season of 2009 suggest that the addition of new members is low, with sporadic immigration pulses as occurs in this, and other areas within the southern Gulf of Mexico (e.g., Martínez-Serrano et al., 2011; Valdés-Arellanes et al., 2011; Morteo et al., 2014, 2017, 2019; Delgado-Estrella, 2015). On the other side, the similarity between the discovery curves of the whole sample, occasional visitors, and transients showed that the dynamics of this population is mainly regulated by the influx of non-resident individuals.

Catchability of RR (mean $0.67 \pm 0.27$ SD) individuals was greater than for OR ($0.35 \pm 0.20$) and OV ($0.32 \pm 0.27$ SD) individuals, and thus abundance estimates for this cluster will be more precise (Williams et al., 2002). In general, catchability should be directly related to research effort alone but, in this case, it probably reflects that regular residents are more catchable by unit of effort than individuals in any other clusters (Williams et al., 2002).

High values of annual survival for the core community (cluster RR + OR) (1.00, 95% CI: 1.00–1.00) contrast sharply to those of occasional visitors: (0.75, 95% CI: 0.59–0.87). As transients were not included in the analysis, and RR and OR clusters behave as a closed population, these highest values for the core community are most probably the result of both a high survival rate and a very low emigration rate.

For long-lived species with complex life history processes, such as cetaceans, adult survival is expected to be high and variable with age, sex, and individual fitness (Ralls et al., 1980). Higher survival rates for resident individuals were indirectly assumed by Morteo et al. (2012b), who showed that these animals interact less frequently with fisheries, when compared to non-residents, thus facing lower exposure to entanglements in fishing gear and retaliation measures by fishers. Therefore, although much lower survival estimates for the visitor fraction is largely based on a high proportion of dolphins permanently leaving the study area shortly after the sampling, it is likely that higher mortality for non-residents is also due to a greater risk of predation by sharks and higher risk of entanglement in fishing gear (Morteo et al., 2012b, 2014, 2017; Reichmont et al., 2018; Morales-Rincón et al., 2019). The latter is supported by reports from recently stranded animals – recorded between 2002 and 2019 (decomposition code 1–3, according to Geraci and Louksbury, 1993) – showing that fisheries-related injuries only occurred in individuals that had not been previously photographed in the study area (Fuentes Del Muro and Morteo, unpublished data, LabMMar-UV).

Survival values recorded for regular and occasional residents in this study are higher than those reported elsewhere (e.g., Wells and Scott, 1990; Speakman et al., 2010; Daura-Jorge et al., 2013; Tezanos-Pinto et al., 2013; Fruet et al., 2015; Vermeulen and Bräger, 2015; Vermeulen et al., 2017; Methion and Díaz López, 2018), probably because of differences in the way we stratified the sample by residency pattern. It is important to notice that the null standard error in the point estimate of survival for both RR and OR is commonly attributed to problems with the data (Cooch et al., 2019, p. 6–24); nevertheless, because of the stratification by residency type, it probably reflects true high values and precision of the estimates due to effective population closure within the study timeframe (i.e., 5-y).

In general, coastal bottlenose dolphins are known to exhibit a wide spectrum of residency patterns, which include transients, seasonal migrants, year-round residents, and a combination of occasional long-range movements, and repeated local residency (Morteo et al., 2014; Wells and Scott, 2018). This seems to be the case for the population in our study area, where Ruiz-Hernández (2014) and Morteo et al. (2019) recorded a limited exchange of individuals between coastal waters off ALS and two northern locations (the Veracruz Reef System and Nautla, see Figure 1), for which dolphins need to travel at least 100 and 230 km, respectively. Our results are similar to other contributions made to the south of our study area, where it was documented that some individuals traveled 270 km on average in a few months, including a dolphin that traveled more than 800 km between the states of Quintana Roo and Tabasco (Delgado-Estrella, 2015, see Figure 1).

Differences in recruitment rate and timing between the core community (RR + OR) and visitors, as found in this study, could correspond to sex-related movement patterns, as reported previously in the Gulf of Mexico, the Caribbean (e.g., Urian et al., 2009; Caballero et al., 2012; Wells, 2014) and also in the study area, through greater residency in females and larger dispersion by males (Morteo et al., 2017, 2019). In this sense, natal site philopatry of both sexes is common and associated to age, as it occurs in the Sarasota population (Wells, 2003, 2014; Sellas et al., 2005). In the northern Gulf of Mexico, molecular data also suggest that some females may move and breed among different communities (Duffield and Wells, 2002), but implications at a local level remain to be investigated.

The existence of two plateaus in the discovery curves of the full sample, OV and OR individuals, indicates at least one occasional pulse of individuals entering from the superpopulation to the study area. These results are consistent with previous findings (Morteo, 2011; Morteo et al., 2014, 2017), suggesting that it is an open population, but the rate of incorporation of new individuals is low in the medium-long term.

Comparable absolute abundances of several estuarine bottlenose dolphin communities around the world are often estimated between 60 and 150 individuals (e.g., Williams et al., 1993; Wilson et al., 1999; Wells, 2003; Balmer et al., 2008; Vermeulen and Cammareri, 2009; Fruet et al., 2011, 2015; Félix et al., 2017). Monthly averages of daily abundances by Morteo et al. (2017) in the study area were about 125 dolphins under the Jolly-Seber model. This number looks consistent with our estimates both within and between years, but results are not directly comparable because of differences in the treatment of the samples.
Our results support the assumption that the dolphin population that uses the coastal waters off ALS is open (Morteo et al., 2017), but emphasizes the existence of a core community of resident individuals that, occasionally, receives an influx of individuals from neighboring waters, with no apparent seasonal trend. It is noteworthy that, unlike other study areas, this core community -living in an open habitat- largely behaves as a closed population. This situation may be more common than previously thought for the species across the Gulf of Mexico, where many individuals remain in relatively small but well provisioned areas, whereas short-term residents and visitors come from the adjacent coastal populations, or have nomadic habits in their constant pursuit of food and mates (Shane, 1980; Irvine et al., 1981; Wells et al., 1987; Ruiz-Hernández, 2014; Delgado-Estrella, 2015; Morteo et al., 2019, among others). The existence of such core communities in the northern Gulf of Mexico has been generally established based on qualitative criteria, mainly the overlap in the presence of transients with year-round and seasonal residents (i.e., Shane, 1980, 1990; Wells et al., 1987; Balmer et al., 2008; Tyson et al., 2011, among others) and, in most cases, refer to dolphin populations living in inshore, estuarine habitats. However, in a coastal habitat adjacent to the Sarasota Bay, Fazioli et al. (2006) found that the dolphin community that prefer the Gulf of Mexico is primarily composed of transients, seasonal residents and individuals with a home range greater than the study area, with fewer year-round residents; nevertheless, details on the process to compute individual residency, and quantitative approaches on the classification of such individuals based on the dynamics of the population is generally lacking.

The permanent presence of dolphin groups and the existence of a core community in the coastal waters off ALS are probably related to a predictable supply of prey and a sheltered environment. Interestingly the dynamic flux of individuals, abundance at a local level (i.e., the core community) seems stable over time, which suggests that a sort of carrying capacity effect is in place. High site fidelity and/or restricted ranging patterns are likely driving population parameters for the core community of bottlenose dolphins in the coastal waters of Alvarado; as this could increase their chances of living in a provisioned habitat, it could also make them more prone to detrimental effects by both documented and currently unknown local threats. Future work should focus on exploring at a finer scale the relationship between these population parameters, the structure of the population (sex/age), and relevant environmental variables for the species.

Differences in residency patterns for the dolphins in our study area could be explained by the socioecological model of Gowans et al. (2007) such that, as resources are spatially and temporally predictable, dolphins remain resident in relatively small areas (Morteo et al., 2014, 2017). Conversely, competition with local dolphins may lead non-resident individuals (mostly males) to range widely to find sufficient resources and mating opportunities (Gowans et al., 2007). This strategy helps to prevent inbreeding (Caballero et al., 2012), and would cause male dolphins of this population to prey on other species that may be available off river mouths and estuaries along coastal waters off the southwestern Gulf of Mexico (Martínez-Serrano et al., 2011; Morteo et al., 2014, 2017, 2019). As the existence of many resident communities has evidenced limited genetic exchange among adjacent sites all along the Gulf of Mexico (Sellas et al., 2005; Caballero et al., 2012; Vollmer and Rosel, 2013), this raises the question of a probable structure at metapopulation level, which should also be investigated.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation. The reduced m-array produced by the MARK output, that contains summary information on the numbers of individuals released at each occasion, and is the basis for the estimation of parameters, is presented in Supplementary Material 3 for reproducibility and replicability.

ETHICS STATEMENT

This animal study was accomplished through federal permits SGPA/DGVS/00351/06, 01407/08 (EM) and SGPA/DGVS/00870/07, 02788/07, 01344/08, and 01649/08 (M. C. Bazúa) issued by the Subsecretaria de Gestion para la Proteccion Ambiental.

AUTHOR CONTRIBUTIONS

JB-J: conceptualization, formal analysis, investigation, methodology, validation, visualization, and writing - original draft. EM: conceptualization, data curation, funding acquisition, methodology, project administration, resources, supervision, validation, and writing - review and editing. CD-A and JB-P: methodology, supervision, validation, and writing - review and editing. PF and ES: conceptualization, methodology, supervision, validation, and writing - review and editing. All authors contributed to the article and approved the submitted version.

ACKNOWLEDGMENTS

This research was part of the first author’s doctoral thesis at Universidad Veracruzana, supported through doctoral fellowship Number 291197/CVU636908 by the Mexican National Council for Science and Technology (CONACyT). This work was also part of the CONACyT project 221750 on the “Trophic ecology of bottlenose dolphins (Tursiops truncatus) and artisanal fisheries interactions in coastal waters off Veracruz State”. Fieldwork was accomplished through federal permits SGPA/DGVS/00351/06, 01407/08 (EM) and SGPA/DGVS/00870/07, 02788/07, 01344/08, and 01649/08 (M. C. Bazúa). Surveys were always conducted by at least one of the authors and many undergraduate students at LabMMar, which were also involved in data collection, photographic analyses and data processing. Israel Huesca helped produce the figures. Local fishers R. Tiburcio, J. Tiburcio and E. Tiburcio always returned the crew safely.
back to shore. Dagmar Fertil and Nathalie Ward reviewed earlier versions of this manuscript. Paul Conn, Manuel Cach, Alberto Delgado Estrella, Tim Gerrodette, Carlos Lira, Manuel Mendoza Carranza, Cecilia Passadore, Grgur Pleslić, Pedro Sánchez Palominio, Kate Sprogis, Gabriela Tezanos Pinto, Fernando Trujillo, and Randy Wells provided valuable information and/or references. This manuscript was greatly improved by comments from MD and AL. This study was part of a collaboration between LabMMar-ICIMAP-IIB, Universidad Veracruzana, and ECOMEGA, Universidade Federal do Rio Grande-FURG. All the fieldwork was conducted in compliance with the “Guidelines for the Treatment of Marine Mammals in Field Research” of the Society for Marine Mammalogy (available at https://marinemammalscience.org/about-us/ethics/marine-mammal-treatment-guidelines).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2021.753484/full#supplementary-material
Residency Shapes Dolphin Population Structure

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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