Sex-Specific Patterns in Abundance, Temporary Emigration and Survival of Indo-Pacific Bottlenose Dolphins (*Tursiops aduncus*) in Coastal and Estuarine Waters

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Inherent difficulties in determining the sex of free-ranging, sexually monomorphic species often prevents a sex-specific focus on estimating abundance, movement patterns and survival rates. This study provides insights into sex-specific population parameters of Indo-Pacific bottlenose dolphins (*Tursiops aduncus*). Systematic, boat-based photo-identification surveys (*n* = 417) were conducted year-round from 2007 to 2013 in coastal and estuarine waters off Bunbury, Western Australia. Pollock's Robust Design was used to quantify population parameters for three datasets: (i) adults and juveniles combined, (ii) adult females and, (iii) adult males. For all datasets, abundance estimates varied seasonally, with general highs during summer and/or autumn, and lows during winter. Dolphins had seasonally structured temporary emigration rates with similar trends between sexes. The derived return rate (1-γ') of temporary emigrants into the study area was highest from winter to spring, indicating that dolphins had a high probability of return into the study area during spring. We suggest that the return of dolphins into the study area and increase in abundance is influenced by the breeding season (summer/autumn). Prey availability is likely a main driver responsible for the movement of dolphins out of the study area during winter. Seasonal apparent survival rates were constant and high (0.98–0.99) for all datasets. High apparent survival rates suggest there is no permanent emigration from the study area. Our sex-specific modeling approach offers a comprehensive interpretation of the population dynamics of a top predator in a coastal and estuarine environment and acts as a model for future sex-based population studies on sexually monomorphic species.

**Keywords:** capture-recapture, capture probability, Robust Design, seasonal movements, sexually monomorphic, Western Australia
INTRODUCTION

Information on population abundance is ecologically important and integral to conservation and management of wildlife (Wilson et al., 1999; Hammond et al., 2002). Population abundance is often estimated through distance sampling (Buckland et al., 2001) and capture-recapture methods (Williams et al., 2002). The advantage of capture-recapture methods is that they allow for the estimation of other important biological parameters, such as temporary emigration and survival (Kendall and Bjorkland, 2001). Capture-recapture models are most informative when integrated with supplementary information on individuals’ sex or age class, thereby allowing for a more holistic interpretation of model outputs (Lebreton et al., 1992; Crespin et al., 2008; Pradel et al., 2008). A sex-based approach to estimating population parameters is important as males and females often vary in their behavioral (social and mating strategies) and ecological (space and habitat use) characteristics (Clutton-Brock, 1989; Whitehead, 1990; van Toor et al., 2011; Sprogis et al., 2016). Hence, sex-based variability in demographic and population parameters can influence population dynamics, population viability, gene-flow, and social behavior (Ruckstuhl and Clutton-Brock, 2005).

Variability between sexes in demographic and population parameters can lead to conservation and management implications, as one sex may be more susceptible to threats than the other (e.g., Crespo et al., 1997; Van Dam et al., 2008; Schofield et al., 2013; Baird et al., 2015). For example, the wandering albatross (Diomedea exulans) exhibits sex-specific differences in foraging areas, which results in a sex-biased human induced mortality (Weimerskirch et al., 1997; Xavier and Croxall, 2005). Specifically, female albatross forage further from the breeding colony than males and, as a consequence, females interact with long-line fisheries more often, resulting in higher levels of bycatch (Xavier et al., 2004). This sex-dependent mortality results in lower survival rates of females, and a subsequent decline in breeding pairs (Xavier and Croxall, 2005). Documentation of whether males and females differ in their abundance, movement patterns and survival rates within a population leads to more comprehensive assessments of potential sex-biased threats and, ultimately, to better-informed conservation measures (Catry et al., 2005; Ruckstuhl and Clutton-Brock, 2005). For many species, however, identifying the sex of free-ranging individuals is often difficult, particularly for sexually monomorphic species (Nichols et al., 2004; Pradel et al., 2008), but also for sexually dimorphic species (Gowans et al., 2000).

Capture-recapture techniques for estimating various population parameters include physical tagging (Baker, 2004), genetic tagging (Palsbøll et al., 1997) or photographic identification (photo-ID) of individuals through their natural markings (Würsig and Jefferson, 1990; Hammond, 2009), including pigmentation patterns (Arzoumanian et al., 2005), pelage marks (Hastings et al., 2008), scars (Gilkson et al., 2007), and whisker patterns (Anderson et al., 2010). Photo-ID has been widely used across taxonomic groups, such as terrestrial and marine mammals (Kelly, 2001; Langtimm et al., 2004), reptiles (Schofield et al., 2008; Gardiner et al., 2014) and fish (Couturier et al., 2014; Kanive et al., 2015). Photo-ID is considered a largely non-invasive method by which animals are initially “captured” in photographs, and subsequently matched through photographic “recaptures” to create an individual’s capture history.

Photo-ID is one of the most commonly used capture-recapture methods for estimating the abundance of cetaceans (Bigg et al., 1990; Hammond, 1990; Würsig and Jefferson, 1990). Natural markings on cetaceans include nicks and notches along the trailing edge of the dorsal fin in bottlenose dolphins (Tursiops spp.; Würsig and Würsig, 1977), color patterns on the ventral surface of the fluke in humpback whales (Megaptera novaeangliae; Katona et al., 1979) and callosity patterns in right whales (Eubalaena australis; Payne, 1986). Photo-ID is most effective when applied to populations with a high proportion of distinctively marked individuals (Hammond, 2009).

Sex-specific population dynamic studies on cetaceans are limited as determining the sex of free ranging individuals is challenging (Morteo et al., 2014; Brown et al., 2016). Some cetacean species are sexually dimorphic in size, but this is not obvious in the field (e.g., the common bottlenose dolphin, Tursiops truncatus, Tolley et al., 1995), or are sexually monomorphic in size, shape and appearance (e.g., the Indo-Pacific bottlenose dolphin, T. aduncus, Hale et al., 2000; Kemper, 2004). The sex of an individual can be documented through several techniques, including molecular sexing (Gilson et al., 1998); laser photogrammetry (Rowe and Dawson, 2009); underwater photography (Webster et al., 2009); opportunistic observations of the genital area (Morteo et al., 2014); or, for females, repeated close association with a calf (Mann, 2000). However, sex determination of a sufficient proportion of individuals within a population to allow for sex-specific capture-recapture analyses requires long-term field effort (Rowe and Dawson, 2009).

Much of our understanding of sex-specific patterns in bottlenose dolphin (Tursiops spp.) social structure and movement ecology arise from long-term studies on T. truncatus in Sarasota Bay, USA (Wells, 1994, 2014), and T. aduncus in Shark Bay, Australia (Connor et al., 2000). Bottlenose dolphins exhibit sex-specific bonds, where their fission-fusion societies favor complex social structuring, typically consisting of mother-calf, adult-male, and mixed-sex juvenile groups (Wells et al., 1987; Smolker et al., 1992). Like most mammals, adult female, and adult male bottlenose dolphins have different strategies to optimize their reproductive output and survival. Adult female dolphins invest care into their offspring and their distribution is influenced by ecological parameters that optimize calf survival, such as the distribution of resources (i.e., prey and protected habitat) and threats (i.e., predators and the mating strategies of males; Connor et al., 2000). Male dolphins do not invest in care of their offspring, and their reproductive output is limited by the number of females they successfully monopolize (Connor et al., 1996). As bottlenose dolphins are polygynous, the distribution of males is heavily driven by the spatial and temporal distribution of receptive females (Emlen and Oring, 1977; Connor and Krützen, 2015). To our knowledge, there are currently no peer-reviewed sex-specific abundance estimates or temporary emigration rates reported for bottlenose dolphins. A sex-specific approach would
provide an opportunity to explore ecological determinants driving any sex-based trends.

A population of *T. aduncus* with seasonal changes in abundance, reside in the coastal and estuarine waters off Bunbury, Western Australia (Smith et al., 2013). There are consistently fewer dolphins present during winter and higher numbers during summer and autumn. The increase in dolphin abundance during summer/autumn coincides with the peak-breeding season, and may be explained by an influx of adult males for breeding purposes (Smith et al., 2013). Here, adult males have larger home ranges than females, suggesting differences in movement patterns between the sexes (Sprogis et al., 2016). The aim of this study was to quantify sex-specific abundance, movement patterns and survival rates of *T. aduncus* across austral seasons to test the hypothesis that there is an influx of males into the study area during summer or autumn months. A sex-specific capture-recapture approach was implemented following Pollock’s Robust Design (herein Robust Design; Pollock, 1982), using photo-ID capture histories.

**METHODS**

**Study Site**

Data were collected year-round from March 2007 to April 2013 off Bunbury (115°63’ E, −33°32’ S), south-western Australia (Figure 1). The Bunbury region typically experiences a Mediterranean climate with cool, wet winters and warm, dry summers. The study site encompassed 120 km², extending 2 km from shore and 50 km along the coast, and consisted of three transect routes: Buffalo Beach, Back Beach and the Inner waters (Figure 1). The Inner waters consisted of the Leschenault Inlet and Estuary, Inner and Outer Harbor, Koombana Bay and the lower reaches of the Collie River (Figure 1). Water depth ranged from <1 m in the estuary to ~15 m offshore.

**Data Collection**

Photo-ID data were obtained during systematic boat-based surveys carried out across austral seasons: summer (December–February), autumn (March–May), winter (June–August), and spring (September–November). A 5 m research vessel with an 80 hp engine was navigated along pre-determined transect routes (Figure 1) at 10 kts. A survey was defined as the traverse of a full transect, with each dolphin group encountered termed a sighting. Surveys were undertaken in weather conditions with Beaufort sea states ≤3, and with two to five observers (median of four). While on transect, observers scanned for dolphins out to approximately 250 m on either side of the vessel.

During each sighting, an experienced researcher with a Nikon D300s camera and a 300 or 400 mm lens aimed to capture a photograph of every dolphin’s dorsal fin for identification purposes (Würsig and Würsig, 1977). For each group encounter, Global Positioning System (GPS) location, time, group composition, and group size were recorded. A group was defined as one or more dolphins within 100 m of any other member involved in the same or similar behavioral activities (Irvine et al., 1981; Wells et al., 1987; Brager, 1999).

Sex of adult dolphins was determined through molecular analyses from biopsy samples collected as part of an associated research project (Manlik et al., unpublished data1), and/or for adult females, repeated, and consistent observations (>three times) in the presence of a dependent calf (Mann et al., 2000). We classified individuals into three age categories: adult, juvenile and calf. Following published work on bottlenose dolphins (e.g., Mann et al., 2000; Ansmann et al., 2013), adults were >2 m (Hale et al., 2000). It is noted that *T. aduncus* in Australia are smaller than other *Tursiops* spp. in other locations and that sizes vary between locations (Connor et al., 2000; Hale et al., 2000). Juveniles were small-bodied compared to adults (approximately two-thirds of adult size) and were not seen consistently beside an adult (Mann et al., 2000). Calves were small (less than two-thirds of adult size) and were consistent with calf length at birth for *Tursiops* spp. in other populations, and maintained repeated close proximity to their mothers (Whitehead and Mann, 2000).

**Data Processing:**

**Photographic-Identification and Photo Grading**

Photographic images of each dorsal fin were used to identify dolphins by unique and permanent nicks and notches (Würsig and Würsig, 1977). Recaptures were matched to a fin catalog using ACDSee 12 software (©2010 ACD Systems International Inc.), following the protocols from the Sarasota Dolphin Research Program (i.e., Bassos-Hull et al., 2013). To help ensure correct identification, matching of “recapture” photos of each dolphin was carried out independently by a minimum of two observers.

Photo-ID data from 2007 to 2009, published in Smith et al. (2013), were incorporated into this study and reanalysed by implementing a photo-grading system, consistent with that applied to the 2010–2013 data. Four independent reviewers graded photographs following the protocols from Rosel et al. (2011; Supplementary Material 1). As such, photographs of each individual documented from 2007 to 2013 were graded for image quality and fin distinguishability in order to minimize heterogeneity in capture probabilities (Stevick et al., 2001; Urian et al., 2015). Image quality affects the likelihood of recognizing an individual, thereby affecting the probability of capture. Fin distinguishability is also important as some individuals are more recognizable than others, which can result in higher probability of capture of more recognizable individuals (Hammond, 1986). Thus, only good and excellent photographs (Q1 and Q2) and highly or moderately distinctive (D1 and D2) dorsal fins were used in analyses (Supplementary Material 1; Rosel et al., 2011). From these images, capture histories of individual dolphins were created and the subsequent capture-recapture analyses were carried out.

**Datasets Used in Analyses**

Analyses were run on three different data sets:

1 Manlik, Oliver, et al., Evolution and Ecology Research Centre, University of New South Wales, Kensington, Australia, unpublished data.
FIGURE 1 | The study site (Bunbury, Western Australia; area = 120 km²) was divided into three transects: Buffalo Beach, Back Beach and the Inner water transect (see insert).

(a) Adults and juveniles combined (including individuals for which sex had not been determined; excluding calves as they were dependent);
(b) Adult females; and
(c) Adult males.

Analyses were also conducted on an additional dataset, which included all known adults in the population, including those of unknown sex (see Supplementary Material 2). In the adult male model, 13 adults were included as male despite not having been confirmed as a male either visually or through molecular analyses. This was justified based on strong and long-term associations with adult males of confirmed sex (see Supplementary Material 3).

Robust Design Structure and Model Assumptions
The Robust Design was implemented to estimate abundance, apparent survival and temporary emigration rates (Pollock, 1982;
Kendall and Nichols, 1995; Kendall et al., 1997). One advantage of the Robust Design is that it accommodates for temporary emigration of individuals, which is not accounted for in closed (does not allow for emigration) or open models (only allows for permanent emigration; Kendall and Nichols, 1995). The Robust Design incorporates both open and closed population models and is structured to have open sampling events (termed “primary periods”), within which are multiple closed events (termed “secondary periods”). Between primary periods, the population is assumed to be open, allowing for births, deaths and permanent or temporary emigration and immigration. Within primary periods, the population is assumed closed, not allowing for change of any kind. In this study, population was defined as the number of individuals frequenting the study area (Williams et al., 2002). Primary periods were based on austral seasons (i.e., four primary periods per year), while the secondary periods were based on the number of days it took to complete the three transects within the study area (Figure 1). Only dolphin sightings that were collected “on-effort” were included in analyses. One sampling regime goal was to complete secondary periods within the shortest time possible (weather dependent) to satisfy the assumption of instantaneous sampling (see below) where possible.

Robust Design capture-recapture models make the following assumptions: (1) marks are unique, permanent and correctly reported; and for all individuals (including unmarked individuals) there are (2) homogenous capture probabilities between individuals within a sampling event; (3) homogeneous capture and recapture probabilities, i.e., no trap response and no heterogeneity; (4) homogenous survival probabilities; (5) instantaneous sampling for secondary periods; (6) the population is closed within primary periods, and (7) captures are independent between individuals (Pollock, 1982; Pollock et al., 1990; Williams et al., 2002).

To minimize violation of the above assumptions, the following steps were taken: (1) to ensure correct identification, unique and permanent nicks and notches were used to identify dolphins, and images were double-checked by two researchers; (2) to reduce heterogeneity in capture probabilities from fin distinctiveness, only graded photographs of good and excellent quality photographs (Q1 and Q2) and distinctive (D1 and D2) dorsal fins were used in analyses (see Supplementary Material 4 for more details on this assumption); (3) capture and recapture was assumed to be homogenous, as animals were not likely to exhibit behavioral responses while photographs were taken because dolphins were not physically captured or handled (see Parra et al., 2006; Nicholson et al., 2012); (4) to ensure homogenous survival probabilities, sex-specific datasets used adults only in analyses, as survival rate can vary by sex (see apparent survival below); (5) to reach instantaneous sampling, secondary periods were completed within the shortest time period possible; (6) to achieve closure, primary periods were structured as seasons rather than years, however this assumption will not be completely satisfied as dolphins in this population temporarily emigrate out of the study area (Smith et al., 2013) and some individuals have larger home ranges than the study area (Sprogis et al., 2016); (7) the model assumption that captures are independent between individuals is most likely violated as dolphin populations are socially driven and the probability of capturing an individual may be increased by capturing its close associates (Connor et al., 2000; Nicholson et al., 2012). This does not bias the estimates, however, but it will inflate the corresponding standard errors (Williams et al., 2002).

Robust Design Analyses

Robust Design analyses were carried out using the program MARK (White and Burnham, 1999). Parameter estimations included: the abundance of animals that are in the study area for each primary period (N), the probability of temporary emigration ($\gamma''$), the probability of apparent survival ($\phi$), the probability of first capture ($r$), and the probability of recapture ($c$).

Temporary emigration is the probability of being temporarily out of the study area if the individual was present ($\gamma''$) or absent ($\gamma'$) in the previous primary period. Three models of temporary emigration were considered. The first was random temporary emigration, where both temporary emigration parameters were equal ($\gamma'' = \gamma'$) and there was no underlying sequential structure of movement (Kendall et al., 1997). The second was Markovian emigration, derived from the Markov process, where the probability of an individual being a temporary emigrant in time $i$ was dependent on whether the individual was present/absent in the study area in time $i-1$ ($\gamma'' \neq \gamma'$; Kendall et al., 1997). Markovian emigration results in a temporally-structured process underlying the movements of individuals (Kendall and Bjorkland, 2001). The influence of temporary emigration for both random and Markovian models was examined as constant over time ($\gamma'' = \gamma'$), time varying over primary periods ($\gamma''$) or varying over austral season (season). Constraints were placed on time-varying ($\phi(t)$) random and Markovian models (i.e., $\gamma''_k = \gamma''_{k-1}$, $\gamma'_k = \gamma'_{k-1}$), so that all parameters could be identified (Kendall, 2011). The random and Markovian models were tested against the third model; the null model of no temporary emigration ($\gamma'' = \gamma' = 0$; Kendall, 2011; Kendall et al., 2012). The return rate ($1 - \gamma'$) of temporary emigrants into the study area from the super-population was also derived. The super-population is defined as the total number of individuals that were present in the study site during the study (Crosbie and Manly, 1985).

Apparent survival ($\phi$) rates represent the product of true survival and permanent emigration (i.e., 1-permanent emigration). Survival probabilities can vary by sex and age class (Baker et al., 2010; Stanton and Mann, 2012). However, it is an assumption of the Robust Design that there is equal probability of apparent survival across all individuals (Burnham et al., 1987; Lebreton et al., 1992). Therefore, the sex-specific adult datasets satisfied this model assumption, as only dolphins that remained in the adult age-class throughout the duration of the study were used. However, the adult and juvenile dataset may have violated this assumption. Seasonal apparent survival rates from primary period $t$ to primary period $t + 1$ were modeled as constant over time ($\phi(t)$) and varying by time ($\phi(t)$).

Capture ($p$) and recapture ($c$) probabilities were assumed to be equal ($p = c$) as photo-ID methods do not require physical handling of animals and, thus, captures are presumed to not affect recaptures (Boyd et al., 2010). Capture probability was modeled as time varying over primary periods ($p = c(t)$), since
environmental conditions were not constant over the duration of the study. Furthermore, capture probability was modeled as constant within primary periods \((p = c(t,))\) or varying by secondary periods within primary periods \((p = c(t, s))\); See Supplementary Material 4 for results and discussion on capture probability.

Overall, a combination of each parameter varying with time, season and constant were used to build models, i.e., 21 different models were run for each dataset: eight time-varying Markovian; seven seasonally dependent Markovian; four random temporary emigration and two no temporary emigration models. The Akaike Information Criterion (AICc) was used for model selection, as this criterion adjusts for small sample sizes. Models with the lowest AICc score were deemed to be most parsimonious (Burnham and Anderson, 2002, 2004).

**Estimating the Proportion of Marked Individuals for Abundance Estimates**

Abundance estimates relate to the marked (D1 and D2) proportion of the population \(\hat{N}_m\). To estimate the total abundance of the population \(\hat{N}_{total}\), estimates were adjusted to account for the proportion of individuals that were unmarked (D3). Sightings in which all individuals were identified from excellent and good quality photographs (Q1 and Q2) were pooled to calculate mark rate \(\hat{\theta}\). Mark rate is the total of marked individuals, divided by the total number of individuals, i.e., including unmarked individuals (excluding calves):

\[
\hat{\theta} = \frac{\text{Total number of marked individuals (D1 and D2)}}{\text{Total number of marked and unmarked individuals}}
\]

Mark rate \(\hat{\theta}\) was not calculated for the sex-specific models, as the sex of all individuals (marked or unmarked) in a group was rarely known. The adult and juvenile combined mark rate was not applied to the adult sex-specific models, as this could diffuse any adult trends. Juveniles are typically not well marked as dolphins accumulate marks with age, hence, a juvenile mark rate would bias the mark rate of adults downwards (Evans and Hammond, 2004; Urian et al., 2015). Therefore, \(\hat{N}_m\) was reported for the sex-specific models, and not \(\hat{N}_{total}\).

For the most parsimonious model for the combined adult and juvenile dataset, mark rate was calculated by year, where \(\hat{\theta}\) was used to adjust the estimated abundance of the population to report \(\hat{N}_{total}\):

\[
\hat{N}_{total} = \frac{\hat{N}_m}{{\hat{\theta}}}
\]

Standard errors for the total population size were calculated using the “delta method” (Seber, 1982; Williams et al., 2002):

\[
SE(\hat{N}_{total}) = \sqrt{\hat{N}_{total}^2 \left( \frac{SE(\hat{N}_m)^2}{\hat{N}_m^2} + \frac{1 - \hat{\theta}}{n \hat{\theta}} \right)}
\]

where \(SE(\hat{N}_m)\) is the standard error of the marked population, and \(n\) is the total number of individuals from which \(\hat{\theta}\) was estimated (i.e., the cumulative number by year). Log-normal 95% confidence intervals were calculated with upper and lower bounds following Burnham et al. (1987):

\[
\hat{N}_{total} = \frac{\hat{N}_{total}}{C} \quad \text{and} \quad \hat{N}_{total}^U = \hat{N}_{total}^L \times C,
\]

where \(C = \exp \left( 1.96 \ln \left( 1 + \frac{SE(\hat{N}_{total})}{\hat{N}_{total}} \right) \right) \)

**RESULTS**

**Sampling Structure and Survey Effort**

In total, 25 primary periods and 139 secondary periods were included in our models (Supplementary Material 5). Secondary periods consisted of 417 surveys (Table 1), including 1310 dolphin group sightings. The time taken to complete secondary periods was weather dependent and ranged from 1 to 21 days \((\bar{x} = 5.14 \pm 0.29 \text{ SE})\). The mean number of days between consecutive secondary periods \((\bar{x} = 11 \pm 1.0 \text{ SE})\) and between primary periods \((\bar{x} = 19 \pm 3.0 \text{ SE})\) varied due to weather dependent data collection.

**Summary Statistics for Model Datasets**

In the combined adult and juvenile dataset, a total of 229 dolphins were highly (D1) or moderately (D2) distinctive. Toward the end of the study, only a few new distinctive dolphins were identified (Figure 2). During each secondary period, the number of dolphins that were identified ranged from 6 to 72 \((\bar{x} = 34.47 \pm 1.13 \text{ SE})\); Supplementary Material 6). The sighting frequency across the duration of the study for individual adult and juvenile dolphins ranged from 1 to 82 \((\bar{x} = 20.92 \pm 1.21 \text{ SE})\; \text{Figure 3} \)

In the adult female dataset, there were 81 highly or moderately distinctive dolphins. Of these, 77 were inferred to be female as they had a dependent calf at some stage throughout the study period. Of these 77 individuals, 36 were also confirmed to be females through molecular analyses. Four females were sexed through molecular analyses only (including one stranded

**TABLE 1 | Summary of annual survey effort covering the three transects, from March 2007 to April 2013.**

| Year | Number of surveys | Number of days | Number of months | Number of dolphin group sightings |
|------|-------------------|----------------|-----------------|----------------------------------|
| 2007 | 29                | 28             | 10              | 114                              |
| 2008 | 73                | 73             | 12              | 235                              |
| 2009 | 69                | 69             | 12              | 242                              |
| 2010 | 81                | 62             | 12              | 204                              |
| 2011 | 72                | 64             | 12              | 217                              |
| 2012 | 69                | 65             | 12              | 225                              |
| 2013 | 24                | 22             | 04              | 73                               |

Total | 417 | 383 | 74 | 1310

*Only on-effort dolphin sightings from completed surveys were included.*
dolphin). During each secondary period, the number of female dolphins identified ranged from 3 to 36 ($\bar{x} = 13.65 \pm 0.48$ SE; Supplementary Material 6). The sighting frequency of adult females ranged from 1 to 82 ($\bar{x} = 23.42 \pm 2.47$ SE; Figure 3) across the duration of the entire study. Across all seasons, adult females were sighted more often than adult males (Table 2).

In the adult male dataset, there were 59 highly (D1) or moderately (D2) distinctive dolphins. Of these, 46 were of known sex based on molecular analyses and 13 through documentation of strong bonds in association (see Supplementary Material 3). The number of male dolphins identified during each secondary period ranged from 0 to 23 ($\bar{x} = 7.77 \pm 0.46$ SE; Supplementary Material 6). The sighting frequency of adult males ranged from 1 to 52 ($\bar{x} = 18.31 \pm 1.75$ SE; Figure 3) across the duration of the entire study.

**Robust Design Abundance Estimates**

The best-fitting model, based on the lowest AICc score, for the combined adult and juvenile dataset was $\phi(.)\gamma(t) \neq \gamma'(t)p = c(t,s)$ with constant survival, time varying Markovian temporary emigration and time varying capture probabilities within primary periods (Table 3; Supplementary Material 7). By contrast, the best-fitting models for the adult female and adult male datasets had seasonally dependent Markovian temporary emigration, $\phi(.)\gamma'(season) \neq c(t,s)$ (Table 3; Supplementary Material 7).

Mark rate for the combined adult and juvenile dataset varied by year, with a low of 0.80 in 2010 and a high of 0.90 in 2008 (Supplementary Material 8). The total number of adults and juveniles combined ($\hat{N}_{total}$) varied by season, with an unprecedented low in winter 2009 (76.23 ± 7.32 SE; 95% CI 67.99–85.48), and high in summer 2010 (184.78 ± 4.46 SE; 95% CI 170.97–199.70; Figure 4; Supplementary Material 8).

The estimated number of marked ($\hat{N}_m$) adult females and males varied between sexes and by season. Abundance estimates for both sexes were, on average, lowest during winter and spring and highest during summer and autumn. For adult females, estimates were consistent among years for spring, apart from an outlier in 2012 with a low of 26.57 ($\pm 1.24$ SE; 95% CI 26.04–33.58). There was also a low in winter 2009 (27.79 ± 2.99 SE; 95% CI 24.97–38.84). The highest estimate was in autumn 2009 (59.56 ± 2.38 SE; 95% CI 57.08–67.70; Figure 5).

For adult males, abundance estimates were quite consistent for autumn; however, they were variable among years for the remaining seasons. Spring had the largest variance in estimated abundance among years, with a low in 2009 ($\bar{x} = 24.02 \pm 5.73$ SE; 95% CI 17.88–43.26) and a high in 2010 ($\bar{x} = 50.16 \pm 5.48$ SE; 95% CI 43.24–66.24; Figure 5).
TABLE 2 | The mean number of sightings ±SE for adult female (n = 81) and adult male (n = 59) dolphins pooled by season.

| Dataset       | Summer       | Autumn       | Winter       | Spring       |
|---------------|--------------|--------------|--------------|--------------|
| Females       | 8.80 ± 0.91  | 6.04 ± 0.60  | 4.23 ± 0.57  | 4.35 ± 0.55  |
| Males         | 7.17 ± 0.86  | 5.10 ± 0.49  | 2.27 ± 0.27  | 3.76 ± 0.41  |

In parentheses is the standardized mean number of sightings per number of secondary periods for summer, autumn, winter, and spring (n = 43, 37, 31, and 28, respectively).

TABLE 3 | The top three Robust Design models, in rank order of AICc scores, for adults and juveniles combined, adult females, and adult males.

| Models                                      | AICc  | ΔAICc | AICc weight | Parameters | Deviance |
|---------------------------------------------|-------|-------|-------------|------------|----------|
| **ADULTS AND JUVENILES COMBINED**           |       |       |             |            |          |
| φ(.) γ*(t) ≠ γ’(t) p = c(t,s)               | 4345.9| 0.0   | 0.999       | 212        | 19724.6  |
| φ(.) γ*(t) ≠ γ’(.) p = c(t,s)               | 4366.3| 20.4  | 0.000       | 190        | 19808.5  |
| φ(.) γ*(season) ≠ γ’(season) p = c(t,s)     | 4366.6| 20.7  | 0.000       | 173        | 19855.9  |
| **ADULT FEMALES**                           |       |       |             |            |          |
| φ(.) γ*(season) ≠ γ’(season) p = c(t,s)     | 3657.1| 0.0   | 0.999       | 172        | 7454.6   |
| φ(.) γ*(.) ≠ γ’(t) p = c(t,s)               | 3681.0| 23.9  | 0.000       | 188        | 7434.2   |
| φ(.) γ*(season) ≠ γ’(.) p = c(t,s)          | 3681.0| 24.0  | 0.000       | 169        | 7486.7   |
| **ADULT MALES**                             |       |       |             |            |          |
| φ(.) γ*(season) ≠ γ’(season) p = c(t,s)     | 2583.5| 0.0   | 0.799       | 173        | 4790.8   |
| φ(.) γ*(season) ≠ γ’(.) p = c(t,s)          | 2586.8| 3.3   | 0.156       | 170        | 4802.2   |
| φ(.) γ*(.) ≠ γ’(.) p = c(t,s)               | 2589.3| 5.9   | 0.044       | 167        | 4812.9   |

The table provides an overview of the Akaike Information Criterion corrected for small sample size (AICc), change in AICc and AICc weight, the number of parameters used in model fit and the deviance explained.

φ apparent survival
γ* ≠ γ’ Markovian temporary emigration.
p = c probability of capture is equal to recapture.
c(t,s) varying by secondary period within primary periods.
(t) time varying, (.) constant, (season) time varying by austral season.

FIGURE 4 | Seasonal abundance estimates \( \hat{N}_{total} \) for adult and juvenile dolphins combined. Lines between data points of the seasonal mean have been used for illustrative purposes only; continuity of values is not implied. Vertical lines show 95% confidence intervals.

Temporary Emigration Patterns
Markovian models performed better than both random temporary emigration and no-movement models (Table 4). Movement for the adult and juvenile dataset had time-varying temporary emigration, whereas adult females and adult males had seasonally dependent temporary emigration. Values for temporary emigration in and out of the study area were similar across datasets. The probability of a dolphin being out of the
study area if the individual was absent ($\gamma'$) in the previous primary period was higher than if the individual was present ($\gamma''$; Table 4). This implied that dolphins temporarily emigrated out of the study area, but subsequently returned.

For $\gamma'$, there was a peak from summer to autumn across all datasets (Table 4). Adult male $\gamma'$ values were lower than adult female values across seasons, apart from summer to autumn. The derived return rate (1-$\gamma'$) of temporary emigrants into the study area from the super-population was highest from winter to spring for all datasets (adult and juvenile = 0.57, adult females = 0.53, adult males = 0.62), indicating that dolphins had a high probability of return into the study area during spring.

For $\gamma''$, there was variability in the lows and highs among datasets (Table 4). For the adult and juvenile population, on average $\gamma''$ was lowest from winter to spring (Table 4). Further, $\gamma''$ had a peak from autumn to winter in 2009 of 0.57 (±0.05 SE), implying that dolphins present in the study area were temporary emigrants during winter, corresponding with an unprecedented decline in abundance (Figure 5). For adult females, $\gamma''$ was generally low from spring to summer (0.099) and high from autumn to winter (0.264). For adult males, $\gamma''$ was low from winter to spring (0.057) and high from spring to summer (0.318). Adult male $\gamma''$ values were lower than those for adult females across seasons, except from spring to summer.

**Survival Rates**

Apparent survival rates were close to 1 at 0.99 (±0.002 SE, 95% CI 0.98–0.99), 0.98 (±0.004 SE, 95% CI 0.97–0.99), and 0.99 (±0.003 SE, 95% CI 0.98–0.10) for adults and juveniles combined, adult females and adult males, respectively. These values strongly suggest there is no permanent emigration in this population and true survival is very high.

**DISCUSSION**

**Abundance Estimates and Temporary Emigration Rates Varied Among Seasons**

We documented seasonal fluctuations in dolphin abundance, with estimates generally lower in winter and higher in summer or autumn. The results derived from datasets using 3 years (2007–2009, Smith et al., 2013) and 6 years (March 2007–March 2013, this study), showed similar seasonal trends with comparable values. Smith et al. (2013) reported abundance estimates for adult and juveniles combined, and reported a low in dolphin abundance during winter 2007 of 63 (95% CI 59–73) and a high in autumn 2009 of 139 (95% CI 134–148). This study, documented a low of adult and juveniles in winter 2009 of 76 (95% CI 68–85) and a high in summer 2010 of 185 (95% CI 171–200). Variations in abundance among seasons and years could be a reflection of both intrinsic (dolphin biology and social dynamics) and extrinsic factors (environmental factors and prey availability).

Seasonal differences in abundance have also been reported in several *T. truncatus* populations. For example, across Florida, there is an increase in abundance during summer in Sarasota (Fazioli et al., 2006), an increase during winter in the Indian River Lagoon (Durden et al., 2011), and an increase in spring and autumn in St. Joseph Bay (Balmer et al., 2008). Conversely, there are no seasonal differences evident for *T. spp.* in Doubtful Sound, New Zealand (Williams et al., 1993), San Diego, USA (Defran and Weller, 1999) and Perth, Western Australia (Chabanne et al., 2012). Thus, the large degree of interpopulation variation in seasonal abundance patterns within species highlights the importance of seasonal sampling.

Temporary emigration for adult and juvenile, adult female and adult male dolphins followed a structured pattern of movement.
(Markovian). These results are consistent with findings by Smith et al. (2013), who reported seasonally structured movement patterns. Furthermore, the probability of an individual dolphin being out of the study area if the individual was absent in the previous primary period was higher than if the individual was present. This implies that dolphins temporarily emigrated out of the study area and subsequently returned. The derived return rate (1−γ′) of temporary emigrants from the super-population was highest from winter to spring, indicating that dolphins had a high probability of return into the study area during spring. Some dolphins were most likely temporary emigrants to the study area (120 km²) due to having larger home ranges (i.e. adult male home ranges reported up to 180 km²; Sprogis et al., 2016) and/or because individual home ranges did not overlap entirely with our study area (Pollock et al., 1990). Therefore, it is likely that several dolphins moved in and out of the study area from the super-population, highlighting the need for temporary emigration to be considered during capture-recapture modeling.

**Biological Factors Affecting Fluctuations in Seasonal Abundance**

The seasonally dependent abundances can be partly explained by the social dynamics of bottlenose dolphins, including their breeding patterns and mating strategies. Bottlenose dolphins have a seasonal breeding pattern, are seasonally polyestrous and have a 12-month gestation period (Connor et al., 2000; Mann et al., 2000). In Bunbury, the peak breeding and calving season is late summer/early autumn (Smith et al., 2016), coinciding with higher dolphin numbers and sighting frequencies, and a return of dolphins into the study area. Compared with females, males form stronger, and more stable bonds (Supplementary Material 3). Such bonds between males are formed to cooperatively gain access to females in order to increase mating success in some populations (Connor and Krützen, 2015). Smith et al. (2013) suggested that the higher abundance of dolphins during summer and autumn may be a result of an influx of adult males for breeding purposes. However, our results do not support this hypothesis. We documented an increase in abundance for adults of both sexes during summer and autumn, with no relative increase in the abundance of males compared to females. There were differences in abundance estimates between sexes that varied across seasons and years, however, confidence intervals were generally overlapping. Furthermore, sighting frequencies of males and females were both relatively high during summer and autumn. However, interestingly, sighting frequencies and capture probabilities for males were lower than for females (see Supplementary Material 4 for discussion on capture probabilities). Therefore, if there was an influx of males, we would have expected to see higher temporary emigration rates of males into the study area compared to females, however, the values were similar for the two sexes. Overall, it is apparent that the higher abundance estimates during summer/autumn are due to an influx of both males and females, which are likely driven by their breeding patterns and prey availability (see below).

**Environmental Factors Affecting Fluctuations in Seasonal Abundance**

Dolphin abundance was generally lower during winter compared to other seasons. Consistent with T. truncatus studies in the USA, similar movements out of study areas occur in winter, most likely in response to fluctuations in prey availability (e.g., Maze and Würsig, 1999; Fazioli et al., 2006; Speakman et al., 2010). Elsewhere, seasonal dolphin movement patterns are also typically linked to prey availability (e.g., Similä et al., 1996; Degrati et al., 2012, 2013). In Bunbury, “potential dolphin prey” in the Leschenault Estuary, Koombana Bay, and near-shore coastal waters has been studied for seasonal abundance, biomass and calorific content (McCluskey et al., accepted). Using three types of fishing gear (seine nets, traps, and gill nets), McCluskey et al. (accepted) documented a decrease in abundance and biomass of prey during winters compared to summers. Additionally, dolphins have been observed >10 km offshore during winter, and not during summer (Sprogis, 2015). Therefore, it is likely that some dolphins temporarily move out of the study area during winter in search of adequate prey resources.

Dolphin movements offshore during winter may be associated with the Leeuwin Current, which flows off the Western Australian coast and transports low salinity waters pole-wards (Godfrey and Ridgway, 1985). There are strong seasonal differences in the Leeuwin Current, in which stronger winter currents flood the continental shelf (Pearce and Phillips, 1988; Cresswell and Griffin, 2004). During this time, primary productivity is enhanced through the entrainment of nutrient rich shelf waters and the formation of eddies (Hanson et al., 2005; Koslow et al., 2008). Ultimately, the strength of the current influences the recruitment and distribution of fish species.
Further investigation into offshore dolphin movements and prey availability are necessary to examine the extent of dolphin movements, and if resource use (e.g., diet and habitat use) differs between sexes (i.e., Rossman et al., 2015).

**Survival Rates Were High and Constant**

The most parsimonious models estimated constant and high apparent survival rates for adult and juveniles combined, adult females and adult males. High survival rates suggest low mortality and no permanent emigration of individuals. Similarly, other *T. aduncus* populations have low permanent emigration (e.g., Nicholson et al., 2012), most likely as a result of natal philopatry (Krützen et al., 2004; Tsai and Mann, 2013). If permanent emigration approaches zero, apparent survival can be represented as true survival. In Bunbury, high survival rates and little permanent emigration are also supported by individual capture histories that spanned the duration of the study.

High and constant survival rates are common for long-lived species with slow growth rates and low fecundity (k-selected species; Gaillard et al., 1998), for example in other cetacean species (e.g., Ramp et al., 2010; Cantor et al., 2012; Tyne et al., 2014). Consistent with this general trend and the findings from our study, relatively high survival rates have also been shown in other *T. aduncus* populations (e.g., Reisinger and Karczmarski, 2010; Mansur et al., 2012; Webster et al., 2014). Our research provides an example for estimating sex-specific survival rates, which is especially important to determine for threatened populations (e.g., Baker et al., 2013) and for populations with sex-biased threats (e.g., Baird et al., 2015).

**CONCLUDING REMARKS**

This study provides insights into sex-specific population parameters of Indo-Pacific bottlenose dolphins in a coastal and estuarine environment. The results highlight that (i) abundance estimates were seasonally dependent and there was an increase of both sexes into the study area during summer and autumn, (ii) temporary emigration rates were seasonal and showed similar patterns between the sexes, (iii) survival was constant and high for both males and females, suggesting little or no permanent emigration from the study area. We suggest the peak breeding season (summer/autumn) motivates the return of dolphins into the study area, influencing the increase in abundance. Prey availability is likely a main reason dolphins move out of the study area during winter/spring (also see McCluskey et al., accepted). Our sex-specific modeling approach allowed for a holistic interpretation of the population dynamics of an apex marine predator in a coastal and estuarine environment.

**ETHICAL STANDARDS**

This study was carried out with approval from the Murdoch University Animal Ethics committee (W2009/06, W2342/10) and was licensed by the Department of Parks and Wildlife (SF005811, SF007986, and SF008624). Biopsy sampling for molecular analyses were carried out as a part of broader study, with data collected in accordance to the Murdoch University Animal Ethics Committee approval (W2076/07; W2307/10; W2342/10), and collected under research permits (SF005997; SF006538; SF007046; SF007596; SF008480; SF009119) licensed by Department of Parks and Wildlife.

**AUTHOR CONTRIBUTIONS**

KS, LB, and KP conceived the study. HR and KS implemented the fieldwork, data organization and data processing. SA collected biopsy samples as a part of a broader study. AK and OM processed biopsy samples and determined the sex of biopsied dolphins. JT assisted with data file processing. KS ran the abundance models and analyses. KS led the writing, with input from LB and KP. All authors critically reviewed the manuscript with final proofs.

**FUNDING**

We thank the funding partners for financial support from the South West Marine Research Program (SWMRP); Bemax Cable Sands, BHP Billiton Worsley Alumina Ltd, the Bunbury Dolphin Discovery Centre, Bunbury Port Authority, City of Bunbury, Cristal Mining, the Western Australian Department of Parks and Wildlife, Iluka, Millard Marine, Naturaliste Charters, Newmont Boddington Gold, South West Development Commission and WA Plantation Resources. KS was supported throughout her PhD by an Australian Postgraduate Award, Murdoch University Research Excellence Scholarship and a SWMRP Scholarship.

**ACKNOWLEDGMENTS**

We thank numerous volunteers and research assistants who helped with fieldwork and data processing. Thank you to our research associates for collecting field data and/or data management; M. Cannon, D. Chabanne, V. Buchanan, K. Nicholson, and B. Goguelat. We are grateful to M. Krützen, D. McElligott, and A. Sellas, who obtained some of the biopsies for genetic analyses. We thank C. Daniel for sexing results from molecular analyses. We are grateful to R. Wells, G. Parra, R. Constantine, B. Würsig, K. Nicholson, and A. Friedlaender and G. Schofield, whose comments greatly improved this manuscript.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmars.2016.00012

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Pollock, K. (1982). A capture-recapture design robust to unequal probability of capture. J. Wildl. Manag. 46, 752–757. doi: 10.2307/3808568

Pollock, K. H., Nichols, J. D., Brownie, C., and Hines, J. E. (1990). Statistical-inference for capture-recapture experiments. Wildl. Monogr. 1–97.

Pradel, R., Maurin-Bernier, L., Gimenez, O., Genovart, M., Choquet, R., and Oro, D. (2008). Estimation of sex-specific survival with uncertainty in sex assessment. Can. J. Stat. Rev. Can. Stat. 36, 29–42. doi: 10.1002/cjs.550360105

Ramp, C., Berube, M., Palsboll, P., Hagen, W., and Sears, R. (2010). Sex-specific survival in the humpback whale Megaptera novaeangliae in the Gulf of St. Lawrence, Canada. Mar. Ecol. Progr. Ser. 400, 267–276. doi: 10.3354/meps08426

Reisinger, R. R., and Karczmarski, L. (2010). Population size estimate of Indo-Pacific bottlenose dolphins in the Algoa Bay region, South Africa. Mar. Mamm. Sci. 26, 86–97. doi: 10.1111/j.1748-7692.2009.00324.x

Ruckstuhl, K. E., and Clutton-Brock, T. H. (2005). “Sexual segregation and the evolution of sex-biased dispersal.” In Sex in Vertebrates, 1st ed. New York, NY; London: Plenum Press), 247–305.

Rucinski, K. E. (2015). Sex-Specific Patterns in Abundance, Home Ranges and Habitat Use of Indo-Pacific Bottlenose Dolphins (Tursiops aduncus) in South-Western Australia. Ph.D, Murdoch University.

Sprogis, K. R. (2015). Sex-Specific Patterns in Abundance, Home Ranges and Habitat Use of Indo-Pacific Bottlenose Dolphins (Tursiops aduncus) in South-Western Australia. Ph.D, Murdoch University.
Würsig, B., and Würsig, M. (1977). The photographic determination of group size, composition, and stability of coastal porpoises (*Tursiops truncatus*). *Science* 198, 755–756. doi: 10.1126/science.198.4318.755

Xavier, J. C., and Croxall, J. P. (2005). “Sexual differences in foraging behaviour and diets: a case study of wandering albatrosses,” in Sexual Segregation in Vertebrates: Ecology of the Two Sexes, eds K. E. Ruckstuhl and P. Neuhaus (New York, NY: Cambridge University Press), 74–91.

Xavier, J. C., Trathan, P. N., Croxall, J. P., Wood, A. G., Podesta, G., and Rodhouse, P. G. (2004). Foraging ecology and interactions with fisheries of wandering albatrosses (*Diomedea exulans*) breeding at South Georgia. *Fish. Oceanogr.* 13, 324–344. doi: 10.1111/j.1365-2419.2004.00298.x

**Conflict of Interest Statement:** 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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