Regional Assessment of the Conservation Status of Snubfin Dolphins (Orcaella heinsohni) in the Kimberley Region, Western Australia

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Implementing conservation measures for data-limited species is a fundamental challenge for wildlife managers and policy-makers, and proves difficult for cryptic marine animals occurring in naturally low numbers across remote seascapes. There is currently scant information on the abundance and habitat preferences of Australian snubfin dolphins (Orcaella heinsohni) throughout much of their geographical range, and especially within the Kimberley region of northern Western Australia. Such knowledge gaps curtail rigorous threat assessments on both local and regional scales. To address this and assist future conservation listings, we built the first comprehensive catalog of snubfin dolphin sightings for the Kimberley. We used these data to estimate the species’ extent of occurrence (EOO) and area of occupancy (AOO) along the region’s 7,000 km coastline, following a simple Bootstrap bivariate kernel approach to combine datasets of varying quality and quantify uncertainty. Our catalog consists of 1,597 visual detections of snubfin dolphins made over a period of 17 years (2004–2020) and collated from multiple sources, including online biodiversity repositories, peer-reviewed scientific articles, citizen science programs, as well as dedicated marine wildlife surveys with local Indigenous communities and Ranger groups. Snubfin dolphins were consistently encountered in shallow waters (<21 m depth) close to (<15 km) freshwater inputs, with high detection rates in known hotspots (e.g., Roebuck Bay, Cygnet Bay) as well as in coastal habitats suspected to be suitable (e.g., Prince Regent River and surrounds, King Sound, Doubtful Bay, Napier Broome Bay and the upper Cambridge Gulf). Bootstrap estimates of EOO and AOO were 38,300 (95% CI: 25,451–42,437) km².
and 700 (656–736) km² respectively, suggesting that snubfin dolphins in the Kimberley are likely Vulnerable under IUCN criteria B2 at a regional scale, in keeping with their global classification. Our study offers insights into the distribution of a vulnerable coastal cetacean species and demonstrates the value of integrating multiple data sources for informing conservation assessments in the face of uncertainty.

**Keywords:** area of occupancy (AOO), extent of occurrence (EOO), citizen science (CS), bootstrap, data integration, geographic range, cetacean marine mammal

### INTRODUCTION

Australian snubfin dolphins (*Orcaella heinsohni*, hereafter ‘snubfins’) are cryptic, small cetaceans endemic to the tropical waters of northern Australia and southern Papua New Guinea. Only differentiated from the closely related Irrawaddy dolphin (*Orcaella brevirostris*) less than two decades ago (Beasley et al., 2005), the species remains poorly known, and is listed as globally Vulnerable under criteria A and C [level VU A2cd + 3cd + 4cd; C2a(i)] of the International Union for the Conservation of Nature (IUCN)’s Red List of Threatened Species (Parra et al., 2017). This classification largely reflects suspected declines in population size stemming from incidental mortality in fishing nets (Parra and Jedensjö, 2014), as well as anticipated habitat loss in the face of growing industrial developments throughout much of nearshore tropical Australia (Allen et al., 2012; Cagnazzi et al., 2013b). A limited body of research indicates that snubfins occur in isolated populations (Brown et al., 2014b) occupying shallow, inshore waters in proximity to freshwater inputs such as estuaries and tidal rivers (Parra et al., 2006b; Palmer et al., 2014b). However, a comprehensive understanding of the spatial ecology, regional distribution, and environmental preferences of snubfins is still lacking.

In particular, while studies of snubfins have been conducted at selected sites in Queensland (Parra, 2006; Parra et al., 2006a,b, 2011; Cagnazzi et al., 2013a,b), the Northern Territory (Palmer et al., 2014a,b), and Western Australia (Thiele, 2010; Brown et al., 2014a, 2016, 2017), extensive portions of the species’ known Australian range have been subject to little or no survey effort (Kaschner et al., 2012). In Western Australia, prominent gaps in sampling coverage remain across the Kimberley, owing to the logistical difficulties and prohibitive costs associated with accessing many parts of the region’s remote, 7,000 km-long coastline (Kordi et al., 2016). Snubfins, named *munumba* in Wunambal and Gaambera languages, have been known to Traditional Owners for several millennia (Karadada et al., 2011), yet the earliest historical account of the species by Europeans within the Kimberley only dates back to 1821 at Prince Regent River (15.452°S; 125.065°E, Figure 1) (King, 1825). Notwithstanding, no targeted sampling was undertaken in the area until 2004, when a limited number of aerial and shipborne visual transect programs were kick-started (Jenner et al., 2008, 2014; Thiele, 2010). While these surveys confirmed the presence of snubfins in low densities within estuaries and coastal embayments, their initial implementation was largely sporadic. Furthermore, observations made in subsequent years by researchers, Indigenous Rangers, and government agencies have never been collated. Such patchy knowledge impedes assessments of the species’ conservation status under IUCN criterion B (“geographic range”) (Allen et al., 2012; Hunt et al., 2017; Smith et al., 2018; Waples and Raudino, 2018), and poses obstacles to its long-term management.

Despite early criticism (e.g., Possingham et al., 2002), there has been increasing appreciation of the utility of IUCN Red Listings (henceforth ‘RLs’) as quantitative instruments for supporting decision-making and conservation priority-setting in the face of such data deficiency (Rodríguez et al., 2006; Hoffmann et al., 2008; Maes et al., 2015; Le Breton et al., 2019). As anthropogenic impacts are not distributed evenly everywhere and management actions are commonly administered at local levels rather than globally (Brito et al., 2010; Pagel et al., 2014; Jenkins and Van Houtan, 2016), several countries around the world have begun to develop their own regional RL schemes aimed at quantifying threats within discrete parts of species’ ranges (Gärdenfors, 2001; Gärdenfors et al., 2008; Nourani et al., 2017). To be rigorous, such efforts ought to be supported by systematic monitoring programs designed around standardized scientific surveys, yet these remain difficult to execute and sustain over the long term in most remote areas (Grech et al., 2014). A promising solution to fill information gaps is to combine different data types from alternative sources, which are often opportunistic in nature (Cheney et al., 2013; Pagel et al., 2014). For instance, non-expert “citizen scientists” now make substantial and active contributions to species mapping by logging ad hoc observations of wildlife using smartphones or other mobile computing technologies (Devictor et al., 2010; Tiago et al., 2017). These can be particularly valuable in biogeographical studies of coastal cetaceans, and can complement or even replace dedicated surveys for tracking distributional changes over broad spatial extents (Embling et al., 2015; Lodi and Tardin, 2018; Alessi et al., 2019). Similarly, community-based approaches that harness the skills, interests, and capacity of Indigenous Ranger organizations are supporting conservation efforts locally, including the monitoring of trends in species and habitats (Grech et al., 2014; Jackson et al., 2015). The latter is particularly relevant where Indigenous people are recognized as land managers in their own right through Native Title, and where they have developed and implemented Healthy Country Management Plans that articulate their vision, objectives and targets for Indigenous Protected Areas (Rist et al., 2019).

In this study, we present the first regional conservation assessment of snubfin dolphins throughout northern Western Australia. Snubfins are currently earmarked as a key target species for fundamental research in Australia (Waples and Raudino, 2018). In 2013, with support from the Australian Federal...
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FIGURE 1 | Visual detections of snubfin dolphins (Orcaella heinsohni) throughout the Kimberley region of Western Australia between 2004 and 2020. Labeled insets display the distribution of both primary and secondary sightings (see Table 1) at key sites, including (A) Roebuck Bay, (B) Cygnet Bay, (C) Prince Regent River, and (D) the Cambridge Gulf. The inset map in the lower right corner shows the boundaries of Indigenous Sea Countries covered by field surveys. These encompass the following Native Title determinations (from north to south): Balanggarra (BAL), Uunguu (UNG), Dambimangari (DMB), Mayala (MYL), Bardi Jawi (BDJ), Bindunbur (BND), and Yawuru (YWR) Sea Countries. Note that Rangers from the Bindunbur community are referred to as Nyul Nyul. Map baselayer: Google © 2019.

Government, efforts were made to develop a National Research Strategy that could help deliver a better understanding of the ecology of tropical inshore dolphins in support of improved conservation decision-making (Department of the Environment, 2013, 2015). Part of this process entailed a technical workshop, during which marine mammal experts were asked to recommend suitable methods for determining the conservation status of snubfins nationally (Brooks et al., 2014). Estimating the species’ extent of occurrence (EOO) and area of occupancy (AOO) under IUCN criterion B was identified as a high priority, on the proviso that such efforts be underpinned by an exhaustive compilation of available datasets, with a strong emphasis on collaboration and Indigenous engagement (Department of the Environment, 2013, 2015). The EOO and AOO are standardized indices of wildlife distribution that are strong and consistent predictors of extinction risk in the marine environment, as larger range sizes provide a buffer against local population declines and mean that species are less likely to experience catastrophic or range-wide losses (Payne and Finnegan, 2007). Importantly, the EOO/AOO are suitable for assessing conservation status even when there is limited information on local threatening processes and the distribution of a taxon is only known from opportunistic observations, unlike other IUCN metrics (Dauby et al., 2017). In particular, criterion B tends to be more suitable than abundance metrics (i.e., IUCN criterion A) for coastal dolphin populations whose sizes are challenging to estimate.
Dolphin sightings (total number, N) were compiled from a variety of sources, including systematic (S) and opportunistic (O) surveys conducted from a range of sampling platforms (vessels, V; kayaks, K; aircraft, A; and land-based sites, L). Surveys undertaken by trained personnel and/or marine mammal experts (primary class) were considered accurate and reliable. Sightings crowdsourced from online repositories or provided by members of the public (secondary class) may be subject to taxonomic reductions as being “observed, estimated, projected, inferred, or suspected”) (IUCN Standards and Petitions Committee, 2000; Guttmacher, 2016; Rueda-Cediel et al., 2018; Fletcher et al., 2019). While this uncertainty is inherently acknowledged in IUCN nomenclature (e.g., criterion A1 specifies population reductions as being “observed, estimated, projected, inferred, or suspected”) (IUCN Standards and Petitions Committee, 2019), it is seldom addressed in practice, with most assessments yielding single threat classifications that do not explicitly reflect the amount and reliability of their underlying inputs (Akçakaya et al., 2000). Some efforts to tackle this issue have been made (e.g., using fuzzy logic or Bayesian networks), but existing solutions are not readily accessible as they rely on proprietary software available only through expensive paywalls (e.g., the RAMAS Red List Pro package) (Akçakaya et al., 2000; Newton, 2010).

Here, we rely on a simple kernel-based weighted resampling approach that treats the veracity of sightings probabilistically, allowing a straightforward assessment of uncertainty. Our analyses build on what is, to our knowledge, the most comprehensive repository of geo-referenced snubfin sightings for the Kimberley. These were obtained from dedicated surveys carried out in partnership with the Indigenous Rangers associated...
with the Yawuru, Dambimangari, Wunambil Gaambera, Bardi Jawi, and Balanggarra Native Title groups, as well as from a range of additional sources such as citizen science initiatives, government archives, and the peer-reviewed literature, among others. Our study provides an example of how to successfully incorporate disparate datasets to conduct RLs of elusive cetaceans in isolated seascapes.

**MATERIALS AND METHODS**

**Study Area**

The study area extends for approximately 7,000 km along the northern coast of Western Australia between Roebuck Bay (122.21°E, 18.17°S) and the Northern Territory border (129°E) (Figure 1). The inshore environment of the Kimberley is a complex matrix of convoluted narrow embayments, drowned valleys, rocky outcrops, and coral-fringed headlands or islands subject to monsoonal climate cycles (Wilson, 2014). Coastal waters are characterized by high levels of turbidity and some of the most extreme tidal ranges in the world, in excess of 10 m in some areas (Wood and Mills, 2008). An array of habitats can be found throughout the region, including mangroves, seagrass meadows, mudflats, and extensive reef systems. These provide foraging, resting, and breeding grounds for a large diversity of marine vertebrates, including cetaceans (Thums et al., 2018).

Supplementary Figure S1 shows the locations of all marine areas, stream outflows, river systems, and reefs/islands referred to throughout the main text. In Western Australia, coastal waters out to three nautical miles fall under the jurisdiction of the State Government, with Indigenous people having management responsibility for their Sea Country where Native Title has been determined (Rist et al., 2019). A network of marine parks has also been implemented across much of the State’s coastal fringe (Supplementary Figure S2), with co-management agreements in place between the State Government and Indigenous people in some cases (e.g., the Uunguu Indigenous Protected Area also encompasses coastal waters). Both marine park and Healthy Country management plans identify marine values within their respective areas and include objectives and targets for their long-term management (Balanggarra Aboriginal Corporation, 2011; Bardi and Jawi Niimidiman Aboriginal Corporation, 2013). Snubfins are recognized as an ecologically important and culturally significant marine species by State and Healthy Country managers across their range.

**Dolphin Data**

**Data Sources**

Sightings of snubfins were collated from a variety of published and unpublished sources and split into two categories reflecting observers’ perceived level of expertise (Table 1). “Primary” sightings consisted of reports made by highly trained personnel (including Kimberley Indigenous Rangers and/or professional marine mammal scientists), and included data from: (a) two 14-day surveys (led by DT) undertaken between Darwin and Broome to describe the presence of snubfins in nearshore tidal river environments (datasets ID OH-001 and OH-002 in Table 1), (b) four 20-day offshore biodiversity surveys conducted under contract with industry (OH-003) in the waters of the Bonaparte Archipelago and Browse Basin (Jenner et al., 2014); (c) 46 daily surveys of Roebuck Bay run along pre-determined transect lines (OH-004) (Thiele, 2010); (d) 36 collaborative surveys undertaken by coastal Indigenous Ranger groups (Nyamba Buru Yawuru, Dambimangari, Balanggarra, and Uunguu) on their respective Sea Countries [OH-005005; see https://bit.ly/snubfin23dolphins-Yawuru for an example (last accessed November 2020)], in addition to a 3-day intensive census of snubfins in the Yawuru Nagulagan/Roebuck Bay Marine Park (Raudino et al., 2019) (Figure 2 and Supplementary Figure S2); (e) extensive dedicated genetic sampling and photo-identification surveys (OH-006) targeting important shallow-water sites along the Kimberley coast, including Beagle Bay, Roebuck Bay, Cygnet Bay, Cone Bay, and the Inner Cambridge Gulf (Brown et al., 2014b, 2016); (f) 14 surveys completed in the northern part of Roebuck Bay in July 2014 (OH-007) concurrently to the deployment of an acoustic logger used for soundscape monitoring (Brown et al., 2017); (h) an array of opportunistic detections reported by biologists and other experienced marine professionals from partner institutions (OH-008 to OH-012) during miscellaneous research and non-research activities, some of which have been published in the peer-reviewed literature (e.g., Allen et al., 2012). Complete or partial GPS tracks were available for most primary datasets and were used to characterize sampling effort (Supplementary Figure S3).

“Secondary” data included incidental sightings collected as part of large-scale aerial surveys targeting dugong (*Dugong dugon*) (OH-013), as well as records crowdsourced from open-access, online biodiversity portals (e.g., Atlas of Living Australia), citizen science mobile applications (DolphinWatch, and Coastal Walkabout), government archives (NatureMap), and anecdotal community reports from members of the public (e.g., recreational ocean users, wildlife tour operators, etc.; OH-014–OH-019, Table 1). With the exception of aerial surveys (Supplementary Figure S3), sampling effort for secondary datasets was unknown (Table 1).

Note that we only considered visual detections, although interest in developing long-term acoustic monitoring programs for the species is rising (Brown et al., 2017; Marley et al., 2017; de Freitas et al., 2018).

**Survey Methods**

A full description of field procedures for individual surveys is given elsewhere (e.g., Brown et al., 2016; Bayliss and Hutton, 2017) and these are only briefly summarized here. On-water surveys were conducted aboard a range of visual platforms, including catamarans, fishing vessels, small-sized rigid inflatables (i.e., zodiacs), research powerboats, and charter yachts (Table 1). All surveys adhered to well-established methodology (Smith and Reeves, 2000; Dawson et al., 2008) with 2–5 experienced observers maintaining constant watch 180° ahead of the platform from the bow, side, and upper bridges (on larger vessels). Visual searches took place during daylight hours, when the boats were motoring or sailing and both weather and visibility conditions were deemed suitable, i.e., Beaufort sea state ≤ 3 (Barlow, 2015).
During systematic surveys, the locations of all dolphin sightings were marked using GPS. Animals were approached at close range whenever possible in order to confirm estimates of group sizes and photograph individual dolphins to support the development of a regional photo-identification catalog⁴.

Aerial surveys were flown aboard a fixed-wing GA8 Airvan in September–October 2015 and May 2017, traveling at 100 knots ground speed and 500 ft, an altitude at which small-bodied marine mammal species can be confidently detected (Bayliss, 1986), though not always differentiated with absolute certainty. Confidence in taxonomic identification typically varies with survey conditions (e.g., Beaufort sea state, glare, water clarity), as well as by observer (e.g., individual experience, eyesight, fatigue levels, and viewing angle from the airplane), and by species (e.g., group size, morphological traits, diving behavior) (Dunshea et al., 2020). Bayliss and Freeland (1989) claimed that after several flights, most observers had little difficulty in telling snubfins apart from other species under favorable viewing conditions. However, other studies reported up to 5–11% of uncertain sightings/disagreements between observers (Hodgson

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⁴https://dolfin.dbca.wa.gov.au/
et al., 2013; Dunshea et al., 2020) (see section “Taxonomic and geographic errors” for the treatment of taxonomic uncertainty).

Surveys followed the procedures outlined in Sobtzick et al. (2013) and Bayliss and Hutton (2017). A team of tandem observers recorded their sightings with at least one experienced observer on each side of the aircraft. Surveys encompassed coastal waters out to the 20 m bathymetric contour between Broome and the Northern Territory border, and were completed in collaboration with Indigenous Rangers following a 5-day intensive training course at Gambimerrri Ranger station on Wunumbal Gaamba Country (Bayliss and Wilcox, 2015).

Data Processing and Treatment

All analyses were undertaken in R v4.0.0. The underlying code is freely available at https://github.com/pboucher/orcaella_eoo.

Data integration is increasingly commonplace in ecological research (Isaac et al., 2020), yet efforts to synthesize vast amounts of heterogeneous information often lead to the assimilation of data burdened with different evidentiary standards (Domisch et al., 2016; Fletcher et al., 2019). Typically, structured datasets obtained during surveys that follow explicit sampling designs are deemed of high quality but remain in short supply, whereas unstructured data collected without strict protocols are growing plentiful but may be ambiguous and subject to important biases (Isaac and Pocock, 2015). Conservation decisions can be highly sensitive to these biases and the potential errors caused by falsely choosing to reject valid observations or accepting anomalous ones (McKelvey et al., 2008), yet many equivocal (but unverifiable) reports are made by experienced amateur naturalists or well-intentioned members of the public whose contributions are difficult to justify excluding (Lee et al., 2014; Brook et al., 2019). The careful treatment of occurrence records is therefore crucial in geographic range assessments of little-studied, rare, and elusive taxa from mixed-certainty datasets (Roberts et al., 2010).

We adopted a two-pronged approach to data processing, focusing firstly on quality control and the correction of a range of common errors (Soberón and Peterson, 2004; Robertson et al., 2016; Zizka et al., 2019; Jin and Yang, 2020), and secondly on the calculation of plausibility (inclusion) scores, which can be interpreted as the probability that each sighting is a true record of the species. The latter step was essential given that snubfins are cryptic and can be easily mistaken for other visually similar and sympatric species (e.g., dugong), especially when detected momentarily from a distance (Bayliss, 1986; Dunshea et al., 2020), and permitted the probabilistic inclusion of all data points, including potentially controversial sightings (Jarić and Roberts, 2014).

Taxonomic and Geographic Errors

Primary sightings, most of which were supported by high-quality digital photographs of the animals (Figures 2A,B), were considered unambiguous and treated as geographically and taxonomically accurate. In contrast, secondary records were likely prone to greater inconsistencies owing to differences in the competence, commitment, and capacity of observers (e.g., eyesight, access to vessels), in addition to the wide range of surveying conditions associated with each data source (Dickinson et al., 2010). Two of the seven secondary datasets (OH-013 and OH-014, Table 1) included a measure of certainty in species identifications, which we relied on to filter out any “guesses,” retaining only those detections labeled as “certain,” “relatively certain,” and “probable.” Sightings from the Atlas of Living Australia (ALA) and the Global Biodiversity Information Facility (GBIF) (OH-018) are subjected to extensive quality tests, and only those that passed taxonomic checks were considered. In the absence of relevant taxonomic information, all other records were assumed to have been correctly assigned to Orcaella heinsohni. Any on-land observations were also randomly reassigned to the nearest ocean area within a 1 km-wide buffer surrounding each point. This allowed us to discard a small number of clearly erroneous sightings (i.e., located several tens to hundreds of km inland), whilst providing a reasonable approximation of the possible distribution of dolphins when the only sighting coordinates available were those of land-based vantage points (e.g., Broome Bird Observatory, Supplementary Figure S4). Both true and likely duplicate records were identified within and across datasets and excluded from the analysis. True duplicates stemmed mostly from the fact that online data infrastructures such as ALA or GBIF often harvest information from common source providers (including each other), and were defined as sightings with identical Julian dates, group sizes, and geographic coordinates (i.e., latitude and longitude) — these were manually discarded. Likely duplicates were taken as sightings made within 100 m of each other on the same date (e.g., repeat observations of the same dolphin group by different citizen scientists). These were spatially thinned using a randomized algorithm, as implemented in the R package spThin (Aiello-Lammens et al., 2015).

Inclusion Probabilities

Recently, several studies have proposed ways of incorporating sighting veracity into ecological inference, either by applying scoring systems developed through expert elicitation (Jarić and Roberts, 2014; Lee et al., 2014, 2015), or via Bayesian frameworks (Solow et al., 2012; Solow and Beet, 2014). The idea of allocating inclusion probabilities to individual sightings is appealing as it relaxes the conventional ‘accept-or-reject’ paradigm of data processing and allows every data point to be fully considered and used, thereby maximizing its analytical value (Brook et al., 2019). However, Bayesian models can be mathematically and computationally intensive, may require prior information that is often missing for data-deficient species, and can be challenging to benchmark against other methods (Saltré et al., 2015). Likewise, expert judgments are necessarily subjective and unlikely to be thoroughly reproducible (i.e., they are prone to vary depending on personal beliefs and experiences, as well as the way in which information is elicited), ultimately limiting their applicability (Martin et al., 2012).

A simpler alternative is to assess ambiguous sightings in terms of their similarity to locations where the species has been documented to occur, for example by down-weighting (or filtering) records associated with environmental conditions that deviate from those encountered in verified datasets collected by
Here, we elected to use the no consensus on how to do so consistently (Gitzen et al., 2006). Specifically, we fitted bivariate kernel density functions to primary sightings within the landscape space defined by both (i) seabed depth and (ii) proximity to the nearest freshwater outflow using the \texttt{bkde2D} function in the \texttt{KernSmooth} package (Wand, 2015). These variables were chosen as the most commonly reported putative drivers of snubfin occurrence throughout their range (Parra et al., 2002, 2006b, 2017; Parra, 2006; Cagnazzi et al., 2013b), and the resulting density surface can thus be thought of as mapping the relative intensity of species presence along two key dimensions of its niche. Depth values were extracted from a high-resolution (30 m) bathymetric grid of the Kimberley curated by Geoscience Australia\(^3\). Freshwater outflows were identified as all perennial streams (major and minor) registered in the Australian Bureau of Meteorology’s Hydrological Geospatial Fabric spatial database v2.1.1\(^4\), which intersected a 1 km buffer around the shore (as marked from the Global, Self-consistent, Hierarchical, High-resolution Geography Database v2.3.7\(^5\)). To account for the complexity of the Kimberley coastline, geodesic distances to freshwater outflows (i.e., shortest ‘least-cost’ trajectories by water, taking into account the presence of intervening land such as islands or capes) (Parra et al., 2006b) were calculated using bespoke code modified from functions available in the \texttt{movecost} R package (Alberiti, 2019). Kernel density estimators are known to be sensitive to the choice of smoothing parameter (i.e., the bandwidth, \(h\)), yet selecting an appropriate bandwidth is notoriously challenging, with currently no consensus on how to do so consistently (Gitzen et al., 2006). Here, we elected to use the \textit{hpi} (‘plug-in’) estimation method implemented in package \texttt{ks} (Duong, 2007). The \textit{hpi} estimator is gaining traction in aquatic research as it has been found to outperform other algorithms and to avoid issues related to convergence failures, skewness, as well as over- and under-smoothing (e.g., Schofield et al., 2013; Jones et al., 2015; Graham et al., 2016; Nifong and Silliman, 2017). Where necessary, \(h\) values were adjusted by visual inspection of the results to ensure that output kernels were in accordance with our prior knowledge about dolphins’ environmental preferences (Sprogis et al., 2016; Doherty et al., 2017). To account for sampling effort, we divided the dolphin kernel by the density surface obtained after applying the same procedure to a set of points placed at regular 10 km intervals along survey track lines (\(N = 3,389\), Supplementary Figure S3; Pennay et al., 2011; Derville et al., 2016). Following the above approach, we set \(h\) to 1 m on the \(y\)-axis (depth) and 1.5 km on the \(x\)-axis (distance to freshwater outflows) for dolphin sightings, and 15 m and 25 km for effort points, respectively. Inclusion probabilities for secondary sightings were taken as the values of the final, standardized kernel (rescaled to 0–1, Brook et al., 2019) at the coordinates of each point in landscape space. All primary sightings were given an inclusion probability of 1 (Figure 3). We also calculated 25 and 90 percent volume contours (PVCs) from the fitted kernel surface. PVCs are slices through the three-dimensional kernel which contain a given percentage of the volume under the fitted surface (e.g., the 90 PVC encloses 90% of the volume beneath the kernel) (Tancell et al., 2012). Doing so allowed us to delineate regions encompassing the densest 25 and 90% of observations (after back-transformation to geographic space), which we interpreted as being indicative of the species’ core areas and full range, respectively (Sveegaard et al., 2011).

### Conservation Assessment

The IUCN relies on two key metrics to assess the conservation status of wildlife species under criterion B: the EOO and the

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\(^3\)http://www.ausseabed.gov.au/surveys-data/data

\(^4\)http://www.bom.gov.au/water/geofabric/

\(^5\)https://www.soest.hawaii.edu/pwessel/gshhg/
AOO (IUCN Standards and Petitions Committee, 2019). The EOO (criterion B1) is defined as the area contained within the outermost boundaries of all known, projected, or inferred records of a species, excluding cases of vagrancy. The AOO (criterion B2) refers to the sub-area occupied by the species within the larger EOO. Both indices are highly relevant to the estimation of extinction risk, as population resilience to catastrophic events may be impaired if either is small or decreasing (Keith et al., 2017). The EOO can be calculated in a number of ways (see Gaston and Fuller, 2009), yet this is routinely done by drawing a minimum convex polygon (MCP) around occurrence points, i.e., the smallest polygon encompassing species records and in which no internal angle is greater than 180°. Despite known caveats (Burgman and Fox, 2003; Borger et al., 2006), IUCN guidelines strongly encourage the use of the MCP for RLs conducted against (Burgman and Fox, 2003; Borger et al., 2006), IUCN guidelines strongly encourage the use of the MCP for RLs conducted against criterion B (IUCN Standards and Petitions Committee, 2019), largely for pragmatic reasons related to conceptual simplicity and the need to estimate EOO consistently across taxa (Joppa et al., 2016). Alternative methods include the α-hull, which is preferred when investigating temporal trends in the EOO, particularly in the context of suspected population declines (see Supplementary Appendix 1). Likewise, multiple techniques have been proposed for measuring the AOO, the most common of which entails superimposing a regular grid over the EOO and summing the area of those cells occupied by the species. The AOO is highly scale-dependent (Hartley and Kunin, 2003; Willis et al., 2003), yet a resolution of 4 km² (i.e., 2 × 2 km cells) is deemed appropriate in most cases, as it ensures that AOO estimates are commensurate with the implicit scale of the thresholds used for delineating categories of threat (i.e., coarser scales preclude listings as critically endangered, whilst finer scales may list more taxa at higher threat categories than the definitions of these categories inherently imply) (Hartley and Kunin, 2003; Willis et al., 2003; Callmander et al., 2007). The AOO can also vary depending on the location of the grid origin (Breiner and Bergamini, 2018), however, this issue can be alleviated by iteratively testing different grid positions and choosing the one that minimizes the number of occupied cells (Moat et al., 2018). Several R packages exist for performing RLs, including Redlistr (Lee et al., 2019), red (Cardoso, 2017), and ConR (Dauby et al., 2017), all with largely similar functionalities. Here, we used ConR to quantify the EOO and the AOO for snubfins across the Kimberley. As per Cagnazzi et al. (2013b), both metrics were computed based on the positions of snubfin groups rather than individuals. We kept the package defaults, which align with IUCN guidelines and treat the EOO as an MCP, computing the AOO at 4 km² resolution (IUCN Standards and Petitions Committee, 2019). For comparative purposes, we also ran a complementary analysis where the EOO was calculated as an α-hull (see Supplementary Appendix 1). We set the nbc.rep.rast.AOO argument to 50, so that 50 raster translations with random starting points would be generated during AOO calculations (Dauby et al., 2017). Importantly, failure to explicitly consider variance in IUCN metrics may lead to incorrect range assessments and inconsistent threat evaluations (Açakayla et al., 2000; Currey et al., 2009). Following Brook et al. (2019), we quantified uncertainty in both the EOO and AOO using a resampling approach, whereby each data point is weighted in proportion to its reliability. In practice, we performed the analyses iteratively on n = 1,000 replicate datasets created by resampling (i) primary sightings with replacement, and (ii) secondary sightings without replacement, but with weights equal to their inclusion probability. We then calculated summary statistics, including means and 95% percentile confidence intervals.

In addition to the EOO/AOO, the focal taxon must fulfill at least two of the three criteria listed under criterion B, namely: (a) populations are severely fragmented or known to exist in no more than a few locations, (b) a continuing decline has been observed, estimated, inferred, or projected in either the EOO, AOO, extent and/or quality of habitat, number of locations or subpopulations, or number of mature individuals, or (c) extreme fluctuations in the above are apparent (IUCN Standards and Petitions Committee, 2019). Recent photo-identification and molecular evidence indicates that snubfins form resident, genetically isolated populations with low migration rates, limited gene flow (Brown et al., 2014b), and potentially high site fidelity (Parra et al., 2006a). Although there is presently no range-wide abundance estimate for snubfins and only scant information on population trends (Parra et al., 2017), no subpopulation studied to date exceeds 250 mature individuals (Parra et al., 2006a; Cagnazzi et al., 2013b; Palmer et al., 2014a; Brown et al., 2016, 2017). Recent projections of the Kimberley marine system to the year 2050 suggest that snubfins are likely to undergo consistent declines in population size as a result of climate warming and other pressures (Boschetti et al., 2020), and it is expected that future changes in abundance will be sufficiently large and pervasive to cause a net reduction for the entire species of at least 20% over a period of two generations (Parra et al., 2017). For these reasons, we assumed that both criteria (a) and (b) were met.

RESULTS

We obtained 3,099 occurrence records of snubfins throughout the Kimberley. After data cleaning and filtering, our final data comprised 1,207 primary and 390 secondary sightings (total N = 1,597) spanning the period 2004–2020 (Table 1). Primary sightings were collected along 34,865 km of vessel-based survey tracklines crossing seven Sea Countries and encompassing a diverse range of habitat types, from mangroves to mud/sand flats, reefs, and seagrass meadows. Sampling effort covered a total depth range of 902 m (median ± SD = 11.5 ± 80.2 m), both in proximity to, and away from, freshwater outflows (median ± SD = 9.7 ± 64.3 km) (Supplementary Figure S3). Consistent with previous findings (e.g., Parra et al., 2006b), the majority of visual detections were made in shallow, nearshore waters less than 21.3 m deep [i.e., N = 1,526 detections within the 95% highest density interval (HDI) of the associated univariate kernel distribution] and no more than 15.4 km from the nearest freshwater outflow (N = 1,533 within the 95% HDI) (Supplementary Figure S5). This pattern held true even after correcting for sampling effort (Figure 3). A small number of primary sightings (N = 11), however, also occurred in deeper waters (up to 39 m) far from any source of freshwater.
FIGURE 4 | Distributions of regional geographic range estimates obtained by weighted resampling for snubfin dolphins (Orcaella heinsohni) in the Kimberley. (A) Extent of occurrence (EOO) calculated as the area of a minimum convex polygon encompassing all occurrence points. (B) Area of occupancy (AOO), measured as the number of 2 x 2 km grid cells occupied by dolphins. Both metrics are expressed in km. A smooth kernel density line is overlaid to facilitate interpretation. Medians and 95% percentile confidence intervals are shown in black as filled circles and segments, respectively.

(up to 63.8 km). Most dolphins occurred in groups of five or less (mean ± SD = 4.4 ± 6.2). The mean inclusion probability (i.e., weights) for secondary sightings was 0.49 (± 0.25 SD), with some marginal variation apparent between datasets (Figure 3). Roebuck Bay (Yawuru Sea Country), Cygnet Bay (Bardi Jawi Sea Country), King Sound (Mayala, Bardi Jawi, Bindunbur Sea Countries), Doubtful Bay (Dambimangari Sea Country), Prince Regent River (and surrounding embayments, Dambimangari Sea Country), the seaward end of Napier Broome Bay (Wunambal Gaambera Sea Country) and the upper parts of the Cambridge Gulf (Balanggarra Sea Country) were identified as areas of high relative probability of snubfin dolphin occurrence (Supplementary Figure S6). The median EOO was 38,300 km² (95% CI = 25,451–42,437 km²) and the median AOO was 700 km² (95% CI = 656–736 km²) (Figure 4). Consequently, the species was identified as “Vulnerable” VU (B2 a,b) at a regional level. Note that range estimates were substantially smaller when the EOO was measured as an α-hull (11,110 km², 95% CI = 11,100–11,120 km²), leading to a potentially higher conservation listing of Endangered (EN) in some Bootstrap iterations (Supplementary Appendix 1).

DISCUSSION

We performed the first cross-cultural geographic range assessment for snubfins in northern Western Australia against IUCN criterion B, based on the largest database of visual sightings collated to date for Orcaella heinsohni in this area. In doing so, we directly addressed an important objective of the National Research Strategy on tropical inshore dolphins (Department of the Environment, 2013, 2015), and tackled several key knowledge gaps currently hindering cetacean conservation (questions 5, 12, and 13 in Parsons et al., 2015). Our results indicate that snubfins may be regionally listed as VU B2(a,b), consistent with the threat category currently assigned to the species globally (Parra et al., 2017). Given that much of the species’ range across the Kimberley lies within jointly managed marine parks or areas managed as Indigenous Protected Areas, this listing highlights the priority need for ongoing monitoring by both the State Government and Traditional Owners. Importantly, our study also confirms snubfins’ strong (though not exclusive) preferences for shallow (<21 m depth) inshore habitats in the vicinity of freshwater outflows (<15 km), in line with previous research (Parra, 2006; Parra et al., 2006b; Palmer et al., 2014b).

Conservation Assessment

Snubfins occur in dynamic nearshore habitats, and are exposed to heterogeneous threats throughout their range (Bejder et al., 2012). In both Queensland and the Northern Territory, strong concerns have been raised about snubfin bycatch in shark control (bather protection) meshing and gillnet fisheries (Soto et al., 2013; Meager and Sumpton, 2016; Tulloch et al., 2020), as well as chronic stress from vessel traffic, noise pollution, and the discharge of contaminants (Cagnazzi et al., 2013a; Palmer and Peterson, 2013) associated with port construction activities, land clearing, and the expansion of aquaculture operations (Cagnazzi et al., 2013b; Parra and Jedensjö, 2014; Munksgaard et al., 2019). By contrast, the Kimberley is renowned for its remoteness and low human population density, but it also has a long history of subsistence, recreational, and commercial fishing (e.g., Molony et al., 2011). Very little is known about or reported on fisheries interactions with snubfins from any of the fishing sectors throughout the Kimberley. Few commercial gillnet fishers operate in the area (only four licenses; DPIRD, 2020), but...
recreational fishing remains a popular activity, and although no incidental entanglements of snubfins have been documented to date (Gaughan and Santoro, 2020; Tulloch et al., 2020), the limited availability of observer programs, combined with ineffective mitigation technologies (Soto et al., 2013) and an absence of records of bycaught animals suggest that under-reporting is likely. Additionally to fishing, the Kimberley also boasts a rapidly growing nature-based tourism industry (Strickland-Munro et al., 2016), produces a large proportion of the country’s offshore/onshore hydrocarbons (Moore et al., 2016), and is affected by heatwaves and other climatic extremes that are growing in frequency and intensity (Le Nohaïc et al., 2017).

These diverse and spatio-temporally varying pressures are likely to have strong impacts on coastal dolphin populations (Wild et al., 2019), especially those following independent evolutionary trajectories (Crain et al., 2009; Wallace et al., 2010). As such, genetically isolated populations of snubfins may be unable to sustain even limited levels of anthropogenic mortality (Meager and Sumpton, 2016). Recent model simulations for Kimberley marine ecosystems indicate that snubfins will also be negatively impacted by future environmental and development pressures (Boschetti et al., 2020), making regional RLs paramount for establishing monitoring baselines for the species going forward (Brown et al., 2014b, 2016, 2017).

The applicability of IUCN criteria at sub-global level has received mounting scrutiny in recent years (e.g., Milner-Gulland et al., 2006; Brito et al., 2010; Maes et al., 2012) in light of disagreements between conservation assessments performed at different scales (Mounce et al., 2018). We are aware of only one other example of a regional RL being conducted for snubfins, which indicated a lower level of threat across the Northern Territory (Palmer et al., 2017). However, both the EOO and AOO can be calculated in multiple ways, including via cartographic (i.e., grid-based) and areographic (i.e., buffer-based) methods, as well as their individual variants (Breiner and Bergamini, 2018) and combinations (Hernández and Navarro, 2007). Different methods will often yield disparate estimates of threat (Gaston and Fuller, 2009), an issue which is exacerbated by the continuing emergence of alternative metrics (e.g., Area of Habitat, AOH; Ocampo-Peñuela et al., 2016; Brooks et al., 2019; Palacio et al., 2020), and by lingering confusion surrounding how the metrics recommended by the IUCN should be implemented in practice (Breiner and Bergamini, 2018). Without a clear consensus on which approach is most appropriate, comparisons of conservation status across areas are likely to be blurred by methodological decisions that remain largely subjective. For instance, ecologists have measured the AOO using grids constructed at a range of spatial resolutions (Gaston and Fuller, 2009; Maes et al., 2012; Groom et al., 2018), from fine (e.g., 1 × 1 km; Fordham et al., 2013) to very coarse (e.g., 100 × 100 km; Scott et al., 2012). Determining a suitable cell size is not trivial and has key implications for characterizing the occupancy-abundance relationship that underpins many conservation and monitoring programs (He and Gaston, 2000; Steenweg et al., 2018). All else being equal, coarse grains risk failing to detect local population declines, while fine grains demand levels of sampling coverage that are typically incompatible with most field budgets (Joseph and Possingham, 2008). Ideally, cell sizes would be selected according to biological relevance, e.g., to match the size of individual home ranges in mobile taxa (Gaston, 1991; Keith et al., 2000). Recent studies suggest that the accuracy of AOO as an indicator of extinction risk may be maximized when grid cells are between 0.1 and 1 times the largest area plausibly impacted by a pressure event (Keith et al., 2017). Under this definition, the optimal resolution for grid-based AOO may be more closely related to the spatial scale and intensity of threats faced by organisms than to their ecology (Keith et al., 2017). Regardless, both ideas pose challenges for data-deficient species like snubfins, for which limited quantitative estimates of habitat use, home ranges or threats currently exist. We note, however, that a number of efforts are underway to map human activities across northern Australian seascapes. Such knowledge is crucial to performing spatial risk assessments and will be key to identifying the proximate factors underlying extinction risk for the species.

Criterion B is also the most commonly misused IUCN criterion (IUCN Standards and Petitions Committee, 2019). Given the aforementioned issues, it is paramount that the precision of EOO/AOO estimates should be considered and reported in RLs (Açıkakaya et al., 2000; Knapp et al., 2003; Wilson et al., 2011; Fourcade et al., 2013; Connors et al., 2014; Rueda-Cediel et al., 2018). Uncertainty accumulates over the assessment process, meaning that the way in which it is handled can profoundly influence classifications of risk (Gillespie et al., 2011) and may ultimately affect management decisions (Regan et al., 2002; D’Eon-Eggerton et al., 2015). This is especially pertinent to RLs undertaken from mixed datasets with potentially ambiguous sightings, whereby multiple uncertainties are likely to be confounded and cannot necessarily be easily disentangled. When this is the case, one of the simplest ways to represent uncertainty is to specify a best estimate and a range of plausible values, as advocated in the Red List guidelines (IUCN Standards and Petitions Committee, 2019). Great strides have been made toward developing strategies for incorporating uncertainty into RLs (Açıkakaya et al., 2000; Knapp et al., 2003; Currey et al., 2009; Newton, 2010), even though this is seldom apparent in the final IUCN listing itself. However, most of these methods are either data-intensive (e.g., Currey et al., 2009), technically advanced (e.g., Bayesian networks; Newton, 2010), or require proprietary software that is not freely available. In this study, we used an intuitive empirical approach to compute confidence bounds around EOO and AOO estimates, which is not predicated on any statistical assumptions and obviates the need for complex modeling machinery. The approach relies on the weighted resampling of the sighting record, and can be easily implemented in the free software package R (Brook et al., 2019) (see code on Github at https://github.com/pjbouchet/orcaella_eoo). Rather than being arbitrarily discarded, ambiguous observations are assigned lower weights (i.e., inclusion probabilities), thereby ensuring that their contribution to the estimation is proportional.
to their veracity. The assignment of inclusion probabilities can be tailored to specific scenarios and need not be subjective if benchmark datasets of high reliability are available, as was the case here. Specifically, we down-weighted snubfin records made in deeper areas away from freshwater stream networks, as these represent conditions in which animals were rarely detected by professional observers (Allouche et al., 2008; Lin et al., 2017; Fletcher et al., 2019) (Figure 3 and Supplementary Figure S5).

**Snubfin Distribution and Habitat Use**

Shallow bathymetry and proximity to freshwater are defining features of the preferred habitats of many dolphin species at cold-temperate latitudes, including Chilean (*Cephalorhynchus eutropia*) (Viddi et al., 2016; Heinrich et al., 2019), Commerson's (*Cephalorhynchus commersonii*) (Garaffo et al., 2011), Hector's (*Cephalorhynchus hectori*), Maui's (*Cephalorhynchus hectori maui*) (Derville et al., 2016), and bottlenose (*Tursiops truncatus*) dolphins (Mendes et al., 2002). In the tropics, Guiana (*Sotalia guianensis*) (Rossi-Santos et al., 2010), snubfin, Irrawaddy (*O. brevirostris*), Indo-Pacific humpback (*Sousa chinensis*), and Australian humpback (*S. sahulensis*) dolphins are also commonly sighted in/around intertidal channels, coastal inlets, creeks, estuaries, and river mouths (Atkins et al., 2004; Parra, 2006; Parra et al., 2006b; Minton et al., 2013; Palmer et al., 2014b; Hanf et al., 2016), although anecdotal evidence of their presence in deeper areas, e.g. around offshore islands several tens of kms from river networks or estuaries also exists (Brown et al., 2017). The drivers of these distribution patterns are not fully understood but likely relate to the availability of prey, as riverine-estuarine systems are typically productive food-rich areas where fish tend to aggregate (Bräger et al., 2003; Travers et al., 2012; McCluskey et al., 2016). Some arguments have been made that higher levels of turbidity at the confluence of fluvial runoffs and tidal fronts/currents may additionally: (1) enhance foraging efficiency, by providing dolphins with visual/acoustic hunting cover (Bräger et al., 2003; Derville et al., 2016), although possibly at the cost of reduced echolocation range (Jensen et al., 2013), and (2) reduce predation risk and increase survival, by concealing dolphins from tiger (*Galeocerdo cuvier*) and bull (*Carcharhinus leucas*) sharks (Wells et al., 1987; Mann, 2000), both of which are regular predators of nearshore dolphins around Australia (Heithaus, 2001; Simpfendorfer et al., 2001; Heithaus and Dill, 2002). To our knowledge, there is presently limited empirical evidence to support either hypothesis. Turbidity fluctuates strongly on daily, monthly, and seasonal scales (Dawe and Collie, 2017), making relationships between dolphin occurrence and water clarity both dynamic and inconsistent across species and regions (Karczmarski et al., 2000; Gomez and Cassini, 2015; Derville et al., 2016). Likewise, patterns of predation risk are complex and geographically variable, with some odontocetes actively avoiding murky habitats (Gannier and Petiau, 2006), while others seem to prefer them (Derville et al., 2016). The prevalence of shark bites on snubfins in the Kimberley is among the highest ever recorded for any dolphin species (Smith et al., 2018), suggesting that snubfins continue to use shallow turbid areas despite high apparent levels of predation risk. In this context, it is thus possible that other forms of interspecific interactions such as antagonistic encounters or resource competition with sympatric species may dictate space use decisions by snubfins to a greater extent than predation. This has been suggested in Queensland, where snubfins may avoid the aggressive and sometimes dominant Australian humpback dolphin males (Parra, 2006).

Within Western Australia, snubfins have been recorded as far south as the Pilbara and the North West Cape/Exmouth Gulf (22.41°S) (Allen et al., 2012; Hanf, 2015). Such sightings remain uncommon, however, and it is unknown whether they reflect infrequent movements by vagrant individuals, the presence of a remnant population residing in the area (Raudino et al., 2018) and/or may signify the potential for a southward distribution shift of the species with warming seas (MacLeod, 2009). By contrast, our results confirm that the coastal fringe of the Kimberley is a stronghold for snubfins, consistent with previous knowledge of dolphin occurrence throughout the region (Supplementary Figure S6). Importantly, we found that the breadth of habitat space utilized by snubfins may be more limited than previously thought, as (1) our best AOO estimate is relatively low (i.e., <1,000 km²), and (2) the 90 PVC represents only 41% of the species’ State-wide range as documented by the IUCN (Parra et al., 2017). Previous studies have shown that numerous marine species tend to occupy relatively small areas within the marginal limits of their range (Strona et al., 2012). This makes strict adherence to IUCN guidelines problematic for species distributed within narrow bands (e.g., along coastlines), as the EOO-MCP is notorious for producing biased estimates of extinction risk (Joppa et al., 2016). Part of the hurdle is that the MCP is a purely spatial metric that does not consider the characteristics necessary to maintain viable populations and fails to exclude patches of unsuitable habitat (de Castro Pena et al., 2014). Because of this, the MCP is susceptible to range overestimation, which can lead to the down-listing of taxa to lower categories of threat (Joppa et al., 2016). By comparison, more flexible methods that allow for discontinuities within a species range may overcome some of these biases (Pelayo-Villamil et al., 2015) and provide more accurate RLs. We abided by IUCN rules to allow comparability with other assessments, yet we consider the results presented in Supplementary Appendix 1 to be more representative of the conservation status of snubfins currently. Again, we emphasize that species cannot be listed under criterion B on the basis of geographic range size alone and must exhibit other symptoms of risk to qualify for threatened status (Collen et al., 2016). Our assessment is therefore predicated on the validity of our assumptions regarding dolphin population size and structure (see section “Conservation assessment”). Further studies are urgently needed to validate these, and elucidate trends in snubfin abundance, connectivity, and home range size.

**Combining Mixed-Certainty Datasets**

We also showed that dolphins are present in areas where they were previously suspected to occur (e.g., Long Reef and Swift Bay, Uunguu Sea Country; King George River, Balanggarra Sea Country; Doubtful Bay and Talbot Bay, Dambimangari Sea
Country) but had never been hitherto documented (Figure 1 and Supplementary Figure S6). Filling such geographic gaps is critical for supporting conservation listings in remote areas (Mounce et al., 2018), and was made possible by pooling information from both conventional and “non-traditional” data sources (sensu Wood et al., 2015) over a period of 17 years. In doing so, we made the implicit assumption that the animals’ distribution throughout the Kimberley has not changed significantly over the last two decades, and we believe this assumption to be reasonable given the current state of knowledge on snubfin ecology and human pressures throughout the Kimberley. Importantly, nearly a quarter of snubfin detections were crowdsourced from reports made by untrained personnel and members of the public (Table 1). Exploiting publicly contributed data from unconventional digital information streams like social media platforms or app-based citizen science programs has been flagged as a key step in improving the current RL framework (Collen et al., 2016). However, opportunistic or anecdotal data are often considered unreliable and have been criticized for compromising assessments of geographic ranges in rare and elusive taxa (McKelvey et al., 2008). Concerns relate primarily to the challenges associated with accounting for many potential — and often unquantifiable — sources of bias, such as imperfect detection, uneven sampling coverage, taxonomic misidentifications, or regional inequalities in resourcing, field capacity, and expertise (Dickinson et al., 2010; Isaac et al., 2014). These biases can undermine the credibility of both global and regional RLs (Mounce et al., 2018), and an ongoing challenge therefore lies in finding appropriate ways of dealing with the largely unstructured data collected outwith the remit of formal scientific studies (Tenan et al., 2016). Probabilistic data integration has accordingly become an active area of methodological innovation in recent years (Bird et al., 2014; Miller et al., 2019; Isaac et al., 2020), but efforts to date have largely focused on the development of integrated predictive models (sensu Isaac et al., 2020), which were outside the scope of the present work. Rather, our objective was to demonstrate a straightforward and practical approach to combining data from disparate sources in the context of RLs, as an alternative to simple data pooling (Fletcher et al., 2019). Key to our approach was the assignment of probabilistic weights to secondary data sources, in proportion to their similarity with the array of environmental conditions that characterize those sightings made by professional observers. In essence, this allowed us to borrow information from multiple vetted primary datasets to deal with the potential weaknesses of secondary datasets, an approach akin to the philosophy behind integrated modeling (Kindsvater et al., 2018). It also obviates the dangers of relying on subjective perceptions about ecological processes to define intrinsic reliability (McKelvey et al., 2008). We note that the strong imbalance in sample sizes between primary and secondary datasets precluded a comparison of their respective contributions to threat categorizations. However, continued data collection and public engagement will likely narrow this gap and allow more formal assessments of their merits and weaknesses, as has been done elsewhere (e.g., van der Velde et al., 2017; Torney et al., 2019).

**CONCLUSION**

Data deficiency remains a pervasive obstacle to wildlife management and conservation globally, and the IUCN is currently unable to assess thousands of organisms because adequate information for those species is lacking (Bland et al., 2015). This issue is exacerbated in tropical environments, where reports of inconspicuous or uncommon taxa are typically sparse and of lesser quality due to cryptic (e.g., boat-shy) animal behavior, uneven sampling effort, as well as the technical difficulties associated with surveying remote and vast seascapes (Braulik et al., 2018). Species that naturally occur in low numbers also make it difficult to assess population trends over time (Brooks et al., 2017). Snubfins are a compelling example of this, as few studies of the animals’ abundance or geographic range have been undertaken since the species was formally described in 2005, except at a few key sites (Parra et al., 2006a,b; Cagnazzi et al., 2013b; Palmer et al., 2014a; Brown et al., 2016). Knowledge gaps such as these limit our capacity to quantify threats and monitor the condition of populations subject to varying pressures (Braulik et al., 2018). Our work illustrates a collaborative, multi-stakeholder approach (Figures 2C–F) to bringing disparate datasets together to build a better picture of the distribution and range of snubfin dolphins within a region regarded as a hotspot of future extinction risk (Davidson et al., 2012). Indigenous Healthy Country management practices and cross-cultural engagement have been critical in informing other IUCN conservation assessments for threatened mammals in Australia and beyond (Balinggarra Aboriginal Corporation, 2011; Bardi and Jawi Niimidiman Aboriginal Corporation, 2013; Ziembicki et al., 2013), and have played a key role in supporting many facets of wildlife management across the Kimberley (Austin et al., 2017, 2019). Such an inclusive and complementary approach built on equal partnerships and mutually beneficial, two-way knowledge exchange should be the norm in areas belonging to and managed by Indigenous people (Woodward et al., 2020). Logical next steps for snubfin research will be to estimate the home range size of individual animals to better approximate their AOO, and to build predictive statistical models that can map dolphin density at fine scales (Syfert et al., 2014). Recent studies are lending growing support to the usefulness of the latter for delimiting the EOO/AAO while simultaneously accounting for the ecological requirements of focal species (e.g., de Castro Pena et al., 2014; Fivaz and Gonseth, 2014; Sousa-Silva et al., 2014; but see Attorre et al., 2013; Breiner et al., 2017 for counter-points); yet while preliminary models have been developed for Australian humpback dolphins in adjacent areas (Haf, 2015), none are yet available for snubfins. The recent gazettel of the Greater Kimberley Marine Park⁴, in combination with the formation of Saltwater Indigenous Protected Areas (Rist et al., 2019) and the recognition of dolphins as a value within them, may provide the impetus necessary to sustain new Indigenous-driven,

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⁴https://parks.dpaw.wa.gov.au/connect/read/great-kimberley-marine-park
collaborative monitoring programs that can generate the data necessary to underpin these efforts.

DATA AVAILABILITY STATEMENT

A video showcasing the collaborative snubfin dolphin surveys undertaken by the Nyamba Buru Yawuru Land and Sea Unit, the Western Australian Department of Biodiversity, Conservation and Attractions, and the World Wildlife Fund Australia on Yawuru Sea Country can be viewed at https://bit.ly/snubfin-dolphins-Yawuru. R code for reproducing the analyses can be found on Github (https://github.com/pjbouchet/orcaella_eoo). All sightings compiled in this study have either been previously published or are in the public domain (see below), with the exception of observations collected by Ranger groups (dataset OH-005, Table 1). Those data are currently supporting ongoing projects and cannot be made readily available at the time of publication. Any requests for information should be directed to the relevant data custodians, namely: Balanggarra Rangers (balanggarra.rangers@klc.org.au), Bardi Jawi Rangers (bardijawirangers@klc.org.au), Dambimangari Rangers (l.vaughan@dambi.org.au), Nyamba Buru Yawuru Rangers (dean.mathews@yawuru.org.au), Nyul Nyul Rangers (nyulnyurlangers@klc.org.au), and Uunguu Rangers (info@wunambalgaambera.org.au). The datasets OH-001, OH-002, OH-004, OH-008, OH-009, OH-011, OH-015, OH-016, OH-017, and OH-019 are available on the Open Science Framework (https://osf.io/kgdu9/). The datasets OH-003, OH-006, OH-007, OH-010 are published in Jenner et al. (2014), Brown et al. (2016), Brown et al. (2017), and Allen et al. (2012), respectively. OH-012 and OH-014 are hosted on NatureMap (https://naturemap.dbca.wa.gov.au/, free registration required). The datasets OH-013 and OH-018 are available from https://data.csiro.au/collections/ and https://ala.org.au/ respectively.

ETHICS STATEMENT

Ethical review and approval was not required for the animal study because no animals were used. All surveys from which sightings were sourced received relevant ethics approvals. Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

AUTHOR CONTRIBUTIONS

DT, PB, SM, KW, HR, FW, Balanggarra Rangers, Bardi Jawi Rangers, Dambimangari Rangers, Nyamba Buru Yawuru Rangers, and Uunguu Rangers contributed to the acquisition of field data and survey designs in various capacities. PB, DT, HR, KW, and SM organized the database and compiled the literature. FW coordinated the participation of Indigenous Ranger groups in dolphin surveys led by DT. PB analyzed the data, in consultation with HR, KW, DT, and SM. PB led the writing of the manuscript, with all authors making active contributions to drafting, critical review, and editorial development. All the authors have approved the final version of the article.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.614852/full#supplementary-material

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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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