



Article (refereed) - postprint

Sequeira, Ana M.M. et al. 2025. Global tracking of marine megafauna space use reveals how to achieve conservation targets.

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Title: Global Tracking of Marine Megafauna Space Use Reveals How to Achieve Conservation Targets

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- 690 Abstract: The recent Kunming-Montreal Global Biodiversity Framework (GBF) sets ambitious goals, but no clear pathway for how zero loss of important biodiversity areas and halting human-induced extinction of threatened species will be achieved. We assembled a multi-taxa tracking dataset (11 million geopositions from 15,845 tracked individuals across 121 species) to provide a global assessment of space use of highly mobile marine megafauna, showing that 63% of the area they cover is used 80% of the time as important migratory corridors or residence areas. The GBF 30% threshold (Target 3) will be insufficient for marine megafauna's effective conservation leaving important areas exposed to major anthropogenic threats. Coupling area protection with mitigation strategies (e.g., fishing regulation, wildlife-traffic separation) will be essential to reach international goals and conserve biodiversity.
- 700 **One-Sentence Summary:** We provide a basis to design a global network of marine protected areas to conserve marine megafauna biodiversity.

Main Text:

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Together with the recently finalised United Nations High Seas Treaty (1, 2), the KunmingMontreal Global Biodiversity Framework (GBF) (3, 4) seeks to protect, conserve and manage at least 30% of oceans. This is a necessary step to support halting the loss of marine biodiversity (GBF Target 3), which has been particularly acute for large marine species (5-7). These include several iconic large marine vertebrates that have been driven to extinction by overexploitation (e.g., the Steller's sea cow – *Hydrodamalis gigas*, the great auk – *Pinguinus impennis*, and the

Japanese sea lion – Zalophus japonicus), and many others currently showing precipitous declines in abundance (e.g., the hawksbill turtle – Eretmochelys imbricata, shortfin mako shark – Isurus oxyrinchus and North Atlantic right whale – Eubalaena glacialis). These mobile and highly migratory marine vertebrates, hereafter marine megafauna, can act as ecosystem and climate sentinels (8; being good surrogates for other biodiversity) and hold key functional roles that
 assist in structuring and maintaining ecosystems (9-11). However, close to a third of species across marine megafauna taxa are now threatened with extinction (5, 12-18).

Certain characteristics of marine megafauna, such as *K*-selected life history traits, place them at priority for systematic conservation planning (i.e., high vulnerability and high irreplaceability; *19*), and make the 'effective conservation' outlined in GBF Target 3 urgently needed. Many also migrate 1000s of km crossing multiple exclusive economic zones (EEZs) and areas beyond national jurisdictions (ABNJ) presenting a challenge for area-based conservation approaches (*20*). Importantly, such approaches are traditionally based on known geographical ranges reflecting historically known boundaries (*18*) or static maps of occurrence (*21*). However,

devising a management plan that effectively conserves migratory species within Ecologically
and Biologically Significant Areas (22) requires an understanding of how the species use space.
Particularly, detecting important marine megafauna areas used for key life-history events, such as breeding or feeding and migratory behaviours, henceforth IMMegAs (to use a term similar to those recognised by IUCN, such as IMMA – Important Marine Mammal Areas or ISRA – Important Shark and Ray Areas) are only tractable using telemetry data (20, 23-27). Despite the challenges associated with collating such data at global scale (28), the detection of global IMMegAs is essential to understand marine megafauna conservation needs to inform global treaties, and should therefore be prioritised for creating the network of marine protected areas

Using telemetry data to understand global space-use by marine megafauna

aimed by GBF (i.e., the planned increase to 30% of area protection).

We assembled a telemetry dataset unparalleled in size and scope (as the result of a global effort initiated by the MegaMove project; 29) by accepting voluntary contributions of tracking data of highly mobile marine vertebrates - here referred to as marine megafauna, despite some (particularly flying birds) being under the 45 Kg threshold (10). Our dataset encompasses over three decades of tracked movements (1985 – 2018) from 15,845 individuals across 121 species, which after curation (30), resulted in 12,794 individual tracks from 111 species, covering 71.7 % of the area of the world's oceans (Fig. 1). Species include flying birds (hereafter birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and manatees), and turtles. See fig. S1 for latitudinal and longitudinal coverage of the dataset, and tables S1-S3, respectively, for lists of species tracked, tracking data details, and species-specific information. According to global assessments by the International Union for the Conservation of Nature (IUCN; 18), of the 111 species considered, ~ 70% have decreasing (54 species) or unknown (23 species) population trends, and more than 50% (58 species) have a threatened conservation status of Critically Endangered (CR), Endangered (EN), or Vulnerable (VU) (table S4).

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Five main regions exhibited the highest effective number of tracked species (as calculated based on the Shannon entropy; 31): the central Indian Ocean, northeast Pacific, Atlantic northeast and northwest, and around Mozambique/South Africa. A few other locations empirically known as having high animal occurrence also showed high number of species (fig. S2). Areas where more tracking data could be made available include southeast Asia, north of Europe (e.g., Spitsbergen and Greenland), Australia, central Pacific Ocean, and western Africa (particularly the southwest Atlantic and Gulf of Guinea) (Fig. 1, fig. S2).

Using properties of the movement detected in the tracking dataset, including speed, direction and movement coherence (30) (fig. S12-S13), we identified IMMegAs based on key behaviours reflected in residency or migratory (including nomadic or dispersive) behaviour. We did this by using an approach (30) able to evaluate these behaviours collectively across multiple tracks without relying on interpolation across highly variable sampling intervals. This is not possible with the traditionally used state-space models that are typically designed to detect behavioural states on single tracks after interpolating position estimates (e.g., 32).

- We then assessed how much of the IMMegAs occurred within existing marine protected areas (MPA, including marine parks; 33) or exclusive economic zones (EEZs; 34) (shown in fig. S3). We used an optimization algorithm to estimate what configuration of the area covered by our tracking dataset would yield the best selection for setting protected areas for marine megafauna, giving priority to grid-cells that are used for both residency and migratory behaviours across multiple taxa (30). For comparison, we repeated this procedure after developing statistical models to predict areas likely to be used for residency or migration for each taxon within the areas covered by our tracking dataset (30). For data used as input for the models see Table 2. After this modelling procedure, we considered the priority grid-cells as those resulting in highest probabilities (i.e., >0.5 and closest to 1) of being an important area across taxa.
- Finally, we assessed the extent to which the GBF's planned increase to 30% in area protection could assist with reducing impacts from marine megafauna's exposure to anthropogenic threats with a global footprint (35), such as fishing (36-38), shipping (39-41), warming (42-45), plastic (46, 47) and noise pollution (48, 49). We identified these as threats based on the IUCN Threats Classification Scheme (TCS) v3.3 (50) complemented with information from existing literature (12, 51-53) and expert knowledge (fig. S4, and see table S4 for details). We then obtained available global threat data for fishing intensity (54), shipping density (55), plastic density (46, 56), and warming (57, 58), and considered noise to be ubiquitous (based on 59) as no noise dataset is currently available at the resolution needed for a global analyses (but see e.g., 60).

Known biases (61-63) associated with uneven sampling and with tagging individuals in known aggregations or colonies were reduced in our analyses as far as possible by using multiple tagging sites for each species and, where applicable, by normalising data to allow for direct comparisons across species and taxa. From specific tests to assess the influence of (i) tagging location bias, (ii) temporal resolution of tracking data (i.e., including only one location per individual per day, in addition to all locations detected), and (iii) spatial resolution (i.e., repeating all procedures at 0.5°, 1° and 2° grid-cells), we found that these potential confounding factors had negligible effects on our main conclusions (fig. S5 – S8). Finally, randomisation of tracks confirmed animals are selectively using space for important behaviours (fig. S14).

795 *Detected ecologically important areas for marine megafauna and extent* 795 *of existing threats*

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We found that, on average, 66.1% of the total area covered by our tracking data was used as migratory corridors (50%) or residencies (44.8%) (Fig. 2A), with ~29% used for both behaviours (30); noting that for sirenians, data were insufficient to detect migratory behaviours (fig. S9). Animals spent on average 90% of their tracked time (estimated using one position per day) within areas where we detected these behaviours (Fig. 2B). Most of this time (~80%) was spent in areas used for residency (or both residency and migration) (fig. S10), with considerable overlap across both behaviours.

On average, only 7.5% of the entire area covered by our tracking dataset occurred inside MPAs (which currently cover ~8% of the global ocean), with ~5% corresponding to areas of detected
 residency or migratory behaviours (Fig. 2). Similarly, animals spent a greater amount of time outside, than inside, MPAs (on average >85%). The time spent inside MPAs corresponded, on average, to 13.6% of all time animals spent displaying residency or migratory behaviours (ranging between 0.3% for polar bears and 23.9% for penguins) (Fig. 2). The results indicate limited opportunity for significant conservation of marine megafauna within the current extent of global MPAs, which were mainly designed to protect specific habitats rather than threatened mobile marine megafauna. However, conservation efforts could be considerably improved in the future by specifically including IMMegAs in new MPA placement.

All space-use and identified residency and migratory behaviours occurred with a ~40-60% split respectively between EEZs and the high seas (which respectively cover 41.3% and 58.7% of the oceans) (Fig. 2). Similar split of space-use between EEZ and high seas was obtained across each taxa, with clear exceptions for sirenians and polar bears (for which most movements occurred inside EEZs). Despite this pattern of space-use slightly biased towards the high seas, most time (on average 74.1%, of which 67.1% corresponded to detected migration or residency) was spent inside, rather than outside, EEZs, and ranged from 61.5% for flying birds to 90.2% for cetaceans (Fig. 2). Although protection of high seas IMMegAs is urgently needed, the large proportion of time animals spend conducting important behaviours within EEZs suggests that an initial focus on enhancing protection within EEZs could provide the fastest benefits for marine megafauna conservation, particularly because implementation may be easier.

To identify what areas could be prioritised for protection, we used an optimisation algorithm (fig. S15 – S16) to select a total of 30% of the 71.7% area covered by our tracking dataset (i.e., 21.3% of the global ocean; Fig. 3). We did this because our tracking dataset does not cover the entire ocean, and also to allow for later additions of new protected areas if other IMMegAs are identified once new tracking data are available. The optimisation algorithm aims to highlight which areas could provide higher representativeness of IMMegAs, but also to indicate where the additional protected areas could be complementary to existing MPAs (sensu 19), which currently fail to represent marine megafauna space-use (25; Fig. 3). Our results show that 30% area protection allows coverage of only less than half of the IMMegAs we discovered (41.6% and 38.8%, respectively, based on data and model predictions; fig. S17), leaving ~60% unprotected (58.4%, and61.2% based on data and model predictions, respectively) (Fig. 3).

835 Our complemented IUCN Threats Classification Scheme(50) (table S4) showed that commercial fishing and climate change affect more than 80% of the species included in our dataset (fig. S4).

Shipping has impacts on species across all taxa, including all turtles, sirenians, polar bears, most species of cetaceans considered, plus five birds, four fishes, five seals, and one penguin. Plastic pollution is a threat for all turtles and seals (but not yet listed on IUCN for leopard seals – *Hydrurga leptonyx*), most cetaceans, and ~35% of birds. Some fishes are also listed as potentially being affected by this threat including two manta rays and five sharks. Noise is listed as affecting all cetaceans, some seals, both sirenians, and also the polar bear, but for the latter this is likely due to potential disturbance of maternal dens on land.

Overlaying the identified (and predicted) areas used by marine megafauna for migration or residency behaviours at a global scale with each of the major global anthropogenic threats considered here (fig. S11), we found that > 96% of IMMegAs are exposed to plastic pollution, shipping and warming, and ~75% to fishing. This exposure includes overlaps within the areas of highest pressure observed for most threats, for example, in the North Atlantic, where we detected important areas for birds, cetaceans, fishes and turtles (Fig. 2 and fig. S9).

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Mitigation strategies will be needed in addition to the proposed increase in area protection to safeguard marine megafauna

Our results reveal that the 30% threshold is insufficient to encompass all IMMegAs globally (Fig. 3), leaving significant conservation risks for marine megafauna. Considering the ubiquity of existing threats, which are pervasive in the IMMegAs we detected (Fig. 3, fig. S11), and the limited scope of the 30% GBF target for area protection, attaining the goal of zero loss of important biodiversity areas and halting human-induced mortality of threatened species seems unlikely (noting some management measures already in place for some species, table S5). Shipping and fishing can in part be alleviated by increasing MPAs (particularly if the highest level of protection is afforded; 64), which can also help reduce noise pollution. However, plastic pollution or climate change impacts will not be alleviated with the planned increase in area protection (even if MPAs can assist improving species resistance and resilience; 65). Therefore, attaining the goal of zero loss of important biodiversity areas will need further action to mitigate anthropogenic pressures.

To reduce exposure of marine megafauna to existing threats and achieve the goals set out in the GBF, the introduction of additional forms of ocean management will be needed, including greater scrutiny of practices and additional direct management decisions with increased enforcement. For example, direct mortality can be reduced by applying fishing thresholds and enforcing standards in fishing operations (including modifications to gear) (*66-70*), and by
developing wildlife-ship traffic separation schemes and slow-down areas (*71, 72*) (e.g., to 2.16 Knots; *73*). If applied in tandem with the increase in protected areas, such interventions will afford marine megafauna a much greater spatial protection from the major threats of industrialised fishing (*23*) and shipping (*41*) known to cause direct mortality (Table 1).

Our analyses show that animals spend the majority of their time within jurisdictions, which presents an opportunity for marine megafauna conservation because individual countries regulate and control most operations within their borders and are therefore able to implement mitigation measures to manage species that use their EEZs. Management of IMMegAs in the high seas, outside national jurisdictions, would benefit from better integration into the United Nations Convention for the Law of the Sea (UNCLOS), and should be considered in the ongoing process to better regulate biological resources in the high seas (*1, 2*). For shipping threats specifically,

International Maritime Organisation regulations can reduce impacts and propel conservation success. For example, the double hull policy resulted in an average reduction of up to 62% in the size of oil spills (74). Engaging (and better regulating) the private sector is another timely way to advance conservation (e.g., 75), as environmental damage is increasingly recognised as a threat to financial stability (75, 76). Past management decisions, either involving the private sector (e.g., end of the whaling industry following the moratorium by the International Convention for Regulation on Whaling; 77) or by listing species on CITES (Convention on International Trade in Endangered Species; 78) have demonstrated success by leading to populations' recovery. However, the drivers of contrasting trajectories of similar populations or species (e.g., right whales increase in the Southern Ocean *versus* decrease in the North Atlantic) are not well understood and likely relate to different exposure to anthropogenic threats.

Creating a larger network of marine protected areas will also greatly benefit from following a systematic conservation planning framework. Although our aim was to identify IMMegAs (rather than outlining what the final 30% of area protection should look like), we followed the 895 initial necessary steps of that framework, including: (i) using marine megafauna biodiversity data (as surrogate for marine biodiversity), (ii) using the set targets from the GBF and UN High Seas Treaty as goal, (iii) focusing on complementing existing MPAs, and (iv) selecting IMMegAs for potential inclusion as MPAs. We then provide a scenario for up to 30% extension of MPAs to show that even if all areas selected specifically included IMMegAs, the 30% protection would still be insufficient to reach set targets, and other mitigation measures will be needed. To follow 900 a systematic conservation planning approach, the final selection of protected areas should also take into consideration aspects not considered here, such as ecosystems of high ecological significance or habitat types that are not yet well represented, as well as considerations of equity and principles of environmental justice (79). It is, however, likely that the final selection of areas 905 for protection will end up being designed to minimise impacts to stakeholders (including the fishing, shipping, energy production and tourism industries). Such possible result further reinforces our conclusion that relying on the 30% area protection will be insufficient to reach the goal of zero loss of important biodiversity areas and halt human-induced mortality of threatened species, and that additional mitigation measures are needed before it is too late.

910 The work we provide here shows the power of assembling tracking datasets to answer pressing conservation concerns. The continued expansion of MegaMove through voluntary contributions will foster greater collaborations allowing to fill data gaps and further reduce biases. Whereas our tracking data covers about 71% of ocean space, the tagging effort was neither random nor uniform in space and time, and 29% of the ocean space was not covered by our dataset

915 (including the central and northwest Pacific ocean). We suggest that statistical models using existing tracking data as input could be used to develop refined global species distributions taking into account animal movements associated with short-term changes in environmental parameters to project the likelihood of encountering animals in areas underexplored by telemetry or bio-logging (80-82).

We also recognise that the available threat distribution data we used here are incomplete and do not include, for example, illegal or artisanal fishing fleets, nor discrimination across fishing gear (which affects species differently). This means that a more detailed spatio-temporal analysis of exposure to threats, as well as an assessment of the vulnerability of different species to specific threats, is required to quantify their potential impacts on species' life-history characteristics.
925 Consideration of the phylogenetic diversity of marine megafauna by examining evolutionary drivers could also be relevant to improve spatial maps. Nevertheless, the IMMegAs we have

identified are key to inform the expansion of existing MPAs to reach the 30% target both within EEZs and in the High Seas.

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Acknowledgments: This research contributes to the MegaMove project endorsed by the UN Decade of Ocean Science (megamove.org). We thank Michelle Heupel for early discussions, Laura Londoño for early assistance with formatting data, and all involved in fieldwork and data 1240 collection - see full details in Supplementary Acknowledgements. We thank Global Fishing Watch and the Global Shark Movement Project for making data available. Ethics and permits information are fully detailed in Supplementary Materials. This study has been conducted using E.U. Copernicus Marine Service Information https://doi.org/10.48670/moi-00021 and https://doi.org/10.48670/moi-00019; NASA Ocean Biology Distributed Active Archive Center (OB.DAAC) data https://oceandata.sci.gsfc.nasa.gov/opendap/SeaWiFS/L3SMI/ and 1245 https://oceandata.sci.gsfc.nasa.gov/opendap/MODISA/L3SMI/contents.html; and the European Centre for Medium-Range Weather Forecasts ERA-Interim Reanalysis product https://www.ecmwf.int/en/forecasts/dataset/ecmwf-reanalysis-interim. Any use of trade, firm, or product names is for descriptive purposes only and does not imply 1250 endorsement by the U.S. Government.

Funding:

1255

A.M.M.S. was funded by a 2020 Fellowship in Marine Conservation by the Pew Charitable Trusts, and ARC DP DP210103091, and through support provided by by the Jock Clough Marine Foundation and three other anonymous donors.

G.C.H. acknowledges funding from the Bertarelli Foundation as part of the Bertarelli Programme in Marine Science (BPMS-2017-4).

D.W.S. acknowledges funding from ERC-2019-ADG 883583 OCEAN DEOXYFISH and NERC NE/R00997X/1.

1260 **V.M.E.** acknowledges funding from MCIN/AEI/10.13039/501100011033 (PID2020-114324GB-C22.

1265	S.A. was supported by Ministerio de Educación, Cultura y Deporte (Spain) [FPU15/01823]. Satellite transmission fees were funded by Universitat Politècnica de València. Tag acquisitions were supported by Universitat Politécnica de València, Asociación Chelonia and Centro de
	Recuperación de Animales Marinos (CRAM).
	(#C314-15), Coordenação de Aperfeiçoamento de Pessoal de Nível Superior - CAPES
0.50	(BJT/A049-2013), and Fundação para a Ciência e Tecnologia - FCT (CEEIND/02566/2021,
1270	UIDP/04292/2020, and LA/P/0069/2020).
	N.P.A.B. was funded by Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) under grant number 482557/2011-7.
	R.D.A. was funded by the North Pacific Research Board and NOAA Fisheries with additional support from Texas A&M University and the Alaska SeaLife Center.
1275	M. Antonopoulou acknowledges seed funding for this work provided by the Emirates Nature – World Wild Fund for Nature office in the UAE and by the numerous private sponsors, listed here

in alphabetical order: 7Days, Abu Dhabi Urban Planning Council, Bridgestone, CASP, College of the North Atlantic, Qatar, Deutsche Bank, Dubai Electricity & Water Authority, Dubai Festival City, Emirates Palace, Environment & Protected Areas Authority, Sharjah, Environment Agency–Abu Dhabi, Fairmont, Géant, Gulftainer, HSBC, Intercontinental, Dubai Festival City, Jebel Ali Golf Resort & Spa, Jumeirah Etihad Towers, Linklaters, Momentum Logistics,

Mubadala, Murjan Marinas, Nokia, Sheikha Salama bint Hamdan Al Nahyan Foundation, The Club, TimeOut Dubai, and the Young Presidents Organisation.
J.A.A. was funded by a scholarship from the Comisión Nacional de Investigación Científica y Tecnológica (CONICYT), Chile, and acknowledges the Chilean Antarctic Institute, which funded the field campaign. GLS tags were provided by the British Antarctic Survey (BAS).
G.A. acknowledges The Rufford Foundation.
I.A. was funded by the Basque Government and the European Data Collection Programme.
M. Auger-Méthé acknowledges the Canadian Research Chairs Program.
R.W.B. acknowledges the US Navy (Pacific Fleet, Atlantic Fleet, Living Marine Resources and Office of Naval Research) and the NMFS (Pacific Islands Fisheries Science Center and Southwest Fisheries Science Center).
A. Barnett acknowledges the Save Our Seas Foundation, Holsworth Wildlife Research Endowment, Winifred Violet Scott Charitable Trust and the Slattery Family Trust.
 E.J. Belda acknowledges that satellite transmission fees were supported by personal research funds made available by the Universitat Politècnica de València. Tags acquisition was funded by Ministerio de Agricultura y Medio Ambiente (16MNSV006); Ministerio de Economía, Industria y Competitividad (CGL2011-30413); Fundación CRAM, Fundación Hombre y Territorio. J. Tomás was supported by project PrometeoII (Generalitat Valenciana) and project INDICIT (EU). Funding for the transmitters came from the JM Kaplan Foundation award to the World Wildlife Fund (WWF) Canada, and from several European institutions: the Spanish International Cooperation Agency (AECI Projects A/2991/05 and A/5641/06), the Spanish Ministry of Education and Sciences (CGL2006-02936-BOS and CGL2011-30413) and the General Foundation of the University of Valencia. The project also received funding from the European Union (Marie Curie Grants FP6 and FP7). A. Bennison was funded by the Irish Research Council (Project ID: GOIPG/2016/503).
 S.R.B. acknowledges financial support and personnel provided by NMFS (Southwest Fisheries Science Center, Southwest Region, Pacific Islands Region, and Office of Protected Resources), and the Tagging of Pacific Predators (TOPP) program of the Census of Marine Life. R.B. received funding from The Wildlife Conservation Society, The Roe Foundation Inc, NABU/Shark Tracker, The National Geographic Society Explorations Council, and The Eppley Foundation. S.E.C. acknowledges funding from the International Governance Strategy of Fisheries and Oceans Canada.
H.A.C. acknowledges Australia Zoo. V.G.C. was funded by the Buenos Aires Zoo, Wildlife Conservation Society, Fondo para la Conservación Ambiental from Banco Galicia, the Cleveland Metropark Zoo–Scott Neotropical

Promoción Científica y Tecnológica FONCyT PICT 2013-2099 Prestamo BID. **R.H.C.** was funded in part by the Dauphin Island Sea Lab, University of South Alabama, Alabama Division of Wildlife and Freshwater Fisheries under traditional Section 6 of the US Fish and Wildlife Service, the Northern Gulf Institute, Mobile Bay National Estuary Program, and the Seamen's Foundation.

Fund, the Inter-American Institute for Global Change Research (IAI) CRN 2076 sponsored by the US National Science Foundation (NSF) grant GEO-0452325, and the Agencia Nacional de

1325 **G.C.** was supported by a Macquarie University Research Excellence Scholarship.

1330

1335

J. Charrassin acknowledges that the Weddell seals tagging study in Dumont d'Urville by LOCEAN laboratory was supported by the Program Terre-Océan-Surface Continentale-Atmosphère from Centre National d'Etudes Spatiales (TOSCA-CNES), and the Australian Animal Tracking and Monitoring System (AATAMS), a facility of Integrated Marine Observing System (IMOS). The Institut Paul Emile Victor (IPEV) programs 109 and 394, Terres Australes et Antarctiques Françaises (TAAF), and the Australian Antarctic Division provided logistical support.

 A. Chiaradia acknowledges funding provided by the Australian Government Research Training Program, Australian Research Council (Linkage Project; LP140100404), Monash University, Phillip Island Nature Parks, Institut Pluridisciplinaire Hubert Curien, ANR-2010-BLAN-1728-01, Australian Conservation Foundation, The Penguin Foundation, Parks Victoria, Centre d'Etudes Biologiques de Chizé and La Rochelle Université, Holsworth Wildlife Research Trust, and Coastcare Australia.

C.C. was supported by the New Caledonian Dugong Technical Committee under the 2010–2015
 Dugong Action Plan in New Caledonia.

E.E.G.C. was supported by the Coral Reef Initiative for the South Pacific (CRISP) mostly funded by the Agence Française de Développement (AFD).

 R.C. acknowledges the New Zealand Ministry for Primary Industries – BRAG; Pew Charitable Trusts; Southern Ocean Research Partnership – International Whaling Commission; Australian Antarctic Division; University of Auckland; Institut de Recherche pour le Développement, France; Conservation International; Blue Planet Marine; Opération Cétacés, New Caledonia; National Marine Mammal Laboratory – NOAA; and the Scientific Committee for Antarctic Research (SCAR), UK.

M.L.C. was funded by a Pew Fellowship Award in Marine Conservation and acknowledges
 European project FEDER Biodiversité.

E. Cuevas-Flores acknowledges the Mexican National Council for Science and Technology -Mexican Ministry of Environment and Natural Resources (project 107770, CAMP-2005-C01-046), the National Fish and Wildlife Foundation (#2006-0091-005), Alliance WWF- Carlos Slim Fund, Chelonia, Inc., Satellite Tracking and Analysis Tool (STAT).

- S. Diamant was supported by the PADI Foundation and IDEA WILD.
 K.L.D. acknowledges funding support from the University of New Hampshire Marine Program, Large Pelagics Research Center, NOAA Grants (#NA04NMF4550391 and #NA10NMF4720028), National Fish and Wildlife Foundation (#2008-0076-000), and the Cape Cod Commercial Fishermen's Alliance.
- 1360 **A.D.M.D.** was funded by the Georgia Aquarium, Darwin Foundation, St Helena Government.

T.K.D. was funded by the UCC Strategic Research Fund.

L.L.D. acknowledges funding provided by the North Carolina Renewable Ocean Energy Program, administered by the Coastal Studies Institute (East Carolina University Outer Banks Campus).

1365 **M.V.E.** warmly thanks the Indonesian Ministry of Environment and Forestry and Ministry of Marine Affairs and Fisheries, the Cenderawasih Bay National Park Authority, the Raja Ampat MPA Management Authority, and the people and government of Raja Ampat, Milne Bay, West

Papua, Bali, East Kalimantan and Nusa Tenggara Timor provinces (especially those from Desa Labuhan Jambu and Desa Kwatisore) for their sponsorship and support, as well the following 1370 donors who financially supported our tagging: the Sunbridge Foundation, SEA Aquarium Singapore, MAC3 Impact Philanthropies, and the Wolcott Henry Foundation, the owners and guests of the MV True North, Asia Coating Enterprise, M.E. Mali, D. Roozen, A. and S. Wong, E. Tan, S. Argyropolous, D. Arnall, A. Hasan, R. Mambrasar, S. Heinrichs, S. Lewis. I. Syakurachman, M. Brooks, P. Rorke-Levy, and S. Neiman. N.E. acknowledges the Dutch Caribbean Nature Alliance. 1375 C.F. acknowledges that sooty tern tracking in Seychelles was funded by the Percy Sladen Foundation, James Cadbury, Robert Gaines-Cooper, Kang Nee, Amanda O'Keefe, Colin & Fiona Short, and Brian & Margaret Jasper.. Funding was provided by DFO (Emerging Fisheries), Government of Nunavut, Nunavut Wildlife 1380 Research Trust Fund, Nunavut General Monitoring Program, Nunavut Wildlife Management Board (#3-09-04), Ocean Tracking Network, University of Windsor, University of Manitoba, ArcticNet Centre of Excellence, Natural Sciences and Engineering Research Council Canadian, Federal Program Office of International, Polar Year (MD-112), Northern Scientific Training Program (Canadian Polar Commission), Polar Continental Shelf Project, W. Garfield Weston Award for Northern Research, and the Molson Foundation. 1385 F.F. acknowledges the Bertarelli Foundation. S.F. acknowledges financial and logistical support provided by Megaptera, its members and friends, the Greenland Institute of Natural Resources, Axa Research Fund, Exagone, Sea Blue Safari, Mikkels Vaerksted, Fondation Nature et Decouverte and the National Geographic Society 1390 Waitt Grant Program. A.S.F. was funded by the Antarctic Wildlife Research Fund (ANT-0823101), NSF Office of Polar Programs (OPP) ANT-0823101, 1250208, and 1440435, IWC, and Southern Ocean Research Partnership awards. C. Garrigue was funded by New Caledonian Government, Ministère de la Transition 1395 Ecologique et Solidaire, WWF for Nature France, Greenpeace International, and Fondation d'Entreprise Total and Opération cétacés NC. **B.J.G.** acknowledges the Natural Environment Research Council (NERC), Darwin Initiative, Marine Turtle Conservation Fund (USFWS, US Department of the Interior). S.D.G. acknowledges the Australian Marine Mammal Centre, Fisheries Research and Development (PN 2005/031), Integrated Marine Observing System (IMOS), DEW, Professional 1400 Association of Diving Instructors (PADI) Project Advancing Wellness and Resilience in Education (AWARE), Australian Bird Environment Foundation, Holdsworth Wildlife Research Endowment, Sea World Research and Rescue Foundation Inc., Nature Foundation of South Australia, South Australian Department for Environment and Heritage Wildlife Conservation Fund, Australian Geographic Society, Norman Wettenhall Foundation, Wildlife Conservation 1405 Fund of South Australia, South Australian Research and Development Institute (SARDI) Women's Bursary 2005, MA Ingram Trust, and Lirabenda Fund (Field Naturalist Society). T.L.G. acknowledges the Save Our Seas Foundation, Swiss Shark Foundation, Watermen Project. Staff and volunteers at Bimini Biological Field Station Foundation. 1410 K.C.H. acknowledges funds provided by NERC and the Department for Business, Energy and Industrial Strategy.

32

N.H. acknowledges support by The Batchelor Foundation, Disney Conservation Fund, Wells Fargo, Guy Harvey Ocean Foundation, Oceana, Oracle, and the West Coast Inland Navigation District.

- 1415 **R.H.** acknowledges funding provided by the Ministry for Business, Innovation and Employment Endeavour Fund C01 1710: "RAMPing-up protection of the Ross Sea", by NZARI (NZ Antarctic Research Institute) and Fisheries New Zealand (respectively), with Regina Eisert as CI, and tags and some field personnel funded by IMOS. The IMOS deployments in Prydz Bay were supported logistically by the Australian Antarctic Division through the Australian Antarctic
- Science Grant Scheme (AAS Projects 2794 & 4329). Work was partially funded by an Australian Research Council Linkage Grant to R.H. and David Slip (LP110200603).
 C.E.H. acknowledges the Earthwatch Institute, David and Lucile Packard Foundation, Wallace Research Foundation, PADI Foundation, and the Arizona-Sonora Desert Museum. Funding was also provided by the Secretaría de Educación y Posgrado Instituto Politécnico Nacional
- 1425 (projects: SIP20141052; SIP20151561; SIP20161935) and the NGO Investigacion, Capacitacion y Soluciones Ambientales y Sociales A.C. **C.E.H.** received a Masters degree bursary from the University of Exeter and the European Social Fund and would like to thank Consejo Nacional de Ciencia y Tecnología (Mexico) for support through a PhD scholarship.
- L.A. Huckstadt was funded under NSF OPP grants ANT-0110687, 0840375, 0533332 and
 0838937, the National Undersea Research Program, and the National Oceanographic Partnership through the Office of Naval Research (ONR).

N.E. Hussey acknowledges funding from the National Sciences and Engineering Research Council of Canada.

C.H. acknowledges support from the Nature Foundation SA Inc., Save Our Seas Foundation,
 Neiser Foundation, Humane Society International, and Mohamed bin Zayed species conservation fund.

R.W.H.I. acknowledges TOPP funding (ONR, NSF, Moore, Sloan, and Packard Foundation) and US EPA GRO fellowship.

- A.A.K. also acknowledges the following management/advisory affiliations/paid consulting
 activities as part of the Top Predator Scientific Working Group of South Africa (Department of Forestry, Fisheries, and the Environment); the Global Shark Movement Project; Shark Spotters NPO (executive committee); South African Whale Disentanglement network (executive committee).
- **D.M.P.J.** acknowledges funding from the National Marine Aquarium (UK) and NationalGeographic.

M.J. was supported by the Science Foundation Ireland Centre for Marine Renewable Energy Research (MaREI 12/RC/2302) and fieldwork supported by the Petroleum Infrastructure Programme (IS13/08) and FishKOSM project funded by the Department of Agriculture, Fisheries and the Marine (15/S/744).

1450 **F.O.L.** acknowledges The Pew Charitable Trusts (PEW) Ocean Science Division and Global Shark Conservation Campaign and Fundação Apolônio Salles de Desenvolvimento Educacional (FADURPE).

1455

P.H.L. thanks CENPES/PETROBRAS (Centro de Pesquisas da PETROBRAS) for supporting the 'Mamíferos e Quelônios Marinhos Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq, PhD scholarship to T.Z.S., process number 141361/2010-7.

J.L. thanks the Founder of the Save Our Seas Foundation for funding and providing all facilities for his work.

G.L. was funded in part by US Fleet Forces Command and managed by Naval Facilities engineering Command Atlantic as part of the US Navy's marine species monitoring program. Additional funding was provided by the National Marine Fisheries Grant to States Program (Grant #NA09NMF4720033) and by private donations managed by the Virginia Aquarium & Marine Science Foundation.

1460

1465

1500

P.L. acknowledges the Italian Consiglio Nazionale delle Ricerche (CNR) and Ministry of Research, University of Pisa, Swedish Natural Science Research Council, NERC, the Darwin Initiative, Italian Space Agency and Accademia Nazionale dei Lincei. The work at Ascension Island was financed by grants from the Swedish Research Council and the Crafoord Foundation to S.Å.

B.C.L.M. acknowledges the Secretaria da Comissão Interministerial para os Recursos do Mar (SECIRM/Brazilian Navy) and CNPq provided grant (#405460/2012-0) for logistics and

1470 equipment through Pró-Arquipélago/Oceanic Islands Program. The Grupo Fundação Boticário de Proteção à Natureza (0760/2007.2), Save Our Seas Foundation (66/2008) and CNPq (478070/2008-0, 482557/2011-7), provided grants for the satellite tags.

A.I.M., **S.D.G.** and **R.H.** acknowledge that satellite tagging of Southern Right Whales in the Great Australian Bight, Australia was funded by a grant from the Department of the

- 1475 Environment Australian Marine Mammal Center, with in kind support from the South Australian Research and Development Institute (Aquatic Sciences), Blue Planet Marine, Macquarie University and Flinders University. They would like to acknowledge the support provided by Dr Mike Double and Dr Virginia Andrews-Goff at the Australian Marine Mammal Centre, and all those who participated in fieldwork.
- M.L. Mallory acknowledges Environment Canada, Natural Resources Canada, Greenland Institute of Nature, Molson Foundation, Natural Sciences and Engineering Research Council of Canada, Canadian Wildlife Federation, and Acadia University.
 J.C.M. acknowledges project funding and equipment provided by NOAA, NMFS Southwest
- Fisheries Science Center, and the National Fish and Wildlife Foundation.
 D.M. acknowledges the BBVA Foundation ('Ayudas Fundación BBVA a Equipos de Investigación Científica 2016'), Spanish Government (grant 'Juan de la Cierva-Formación' FJCI-2014-20064), Fundación Reina Sofía (LIBERA 2017), and NOAA.

A.M. acknowledges the Doñana Biological Station (EBD-CSIC), Consejería de Medio Ambiente y Ordenación del Territorio (CMAOT) of Junta de Andalucía, the Andalusian Marine Environment Management Center (CEGMA) and the NGO Equinac.

Environment Management Center (CEGMA) and the NGO Equinac.
 M. Marcoux was funded by the DFO Nunavut Implementation Fund and the Strategic Program for Ecosystem-Based Research and Advice. Narwhal tagging efforts were supported from the World Wildlife Fund and the Nunavut Wildlife Management Board.

L.M. was principally supported through the Australian Government's Fisheries Research and Development Corporation (FRDC) Grants Scheme (PN 2005/031), co-funded by the South Australian Sardine Fishery. We also thank the Nature Foundation South Australia for financial assistance that supported the purchase of GPS units.

G.M. was principally supported by Xunta de Galicia, Spain, throught Isabel Barreto Programme (2009–2012) and FCT grants (PTDC/MARBIO/4458/2012; IF/01611/2013; NORTE-01-0145-FEDER-000031).

M.M.C.M. acknowledges the MEOP-BR (Marine Mammals Exploring the Oceans Pole to Pole; Brazil), an International Polar Year (IPY) programme funded by the Brazilian Science, Technology and Innovation Ministry through the Brazilian National Research Council (CNPq, Grant no. 520196/2006-6).

M.A.C.N. acknowledges the petrel tracking programme and Malcolm Nicoll were supported by NERC (Grant NE/H5081500) with in-situ support from MWF and NPCS.
 W.J.N. was supported by a Fulbright Fellowship and a Marshall Fellowship during the period field research in Baja California was conducted.

B.M.N. and S.D.R. acknowledges the many supporters, funders, donors and volunteers ofECOCEAN Inc.

1515

B.P. received a Commonwealth Scientific and Industrial Research Organisation (CSIRO) Marine Research scholarship.

E.O. acknowledges funding provided by NOAA NMFS, the Institute of Marine Research in Norway, the Nordic Council of Ministers, and FCT (grants: SFRH/BD/32520/2006 and SFRH/BPD/29841/2006).

S. Oppel acknowledges that work on Ascension Island was partly funded by a Darwin Grant (# 19026) to Ascension Island Government and the University of Exeter (**A.C.B.** and **B.J.G.**), managed on-island by **N.W.** and **S.B.W.**. The king eider study was funded by the Coastal Marine Institute (University of Alaska, Fairbanks), Minerals Management Service, US Geological

- 1520 Survey (Outer Continental Shelf Program), and Canadian Wildlife Service. Further financial and technical support was provided by the Sea Duck Joint Venture, USFWS, North Slope Borough, ConocoPhillips Alaska Inc., Inuvialuit Wildlife Management Advisory Council, WWF, BP Exploration Alaska, Polar Continental Shelf Project, US Geological Survey Alaska Cooperative Fish and Wildlife Research Unit, Institute of Arctic Biology (University of Alaska Fairbanks),
- 1525 and German Academic Exchange Service. The work on St Helena was partly funded by Enterprise St Helena and the Seabird Group. The David and Lucile Packard Foundation, Darwin Plus: Overseas Territories Environment and Climate Fund, the Sir Peter Scott Commemorative Expedition to the Pitcairn Islands, generous donors, and the Royal Society for the Protection of Birds (RSPB) helped to fund our research.
- 1530 **A.M.P.** acknowledges the U.S. Geological Survey (USGS) Ecosystems and Climate and Land Use Change Mission Areas, the USGS Changing Arctic Ecosystems Initiative, and the Bureau of Land Management for primary funding. Additional support was provided through a National Science Foundation grant to the University of Wyoming; the United States Fish and Wildlife Service, Marine Mammals Management and the Arctic National Wildlife Refuge; Environment
- and Climate Change Canada; the North Slope Borough, Department of Wildlife Management;
 the Polar Continental Shelf Project; Polar Bears International; the University of California, Santa Cruz; the San Diego Zoo Wildlife Alliance; and the University of Alberta.
 V H P. asknowledges EU INTERPEC project FAME: The Enture of the Atlantic Marine.

 V.H.P. acknowledges EU INTERREG project FAME: The Future of the Atlantic Marine Environment (2009-1/089) and by LIFE+Berlenga (LIFE13 NAT/PT/000458). Strategic program of Marine and Environmental Sciences Centre (MARE), financed by FCT (MARE –

1540 of Marine and Environmental Sciences Centre (MARE), financed by FCT (MARE – UID/MAR/04292/2013). LIFE Project Marine Important Bird Areas (2004–2008) by the EU INTERREG Project FAME: The Future of the Atlantic Environment (2010–2012) founded by the EU.

D.M. Palacios and **B.M.** acknowledge support provided by the TOPP program of the Census of Marine Life, the US Minerals Management Service, the US ONR (Grants 9610608, 0010085,
1550	0310861, N0014-02-1-0885, N0-176A, and N00014- 09-1-0453), NSF, the Alfred P. Sloan Foundation, the Moore Foundation, the Packard Foundation, the National Geographic Society, the IWC (with funds provided by Exxon Neftegas Limited and Sakhalin Energy Investment Company), the National Research Foundation of South Africa (GUN number 2047517), NOAA through the Northeast Consortium, based at the University of New Hampshire (Grant #NA16FL1324), the International Association of Oil and Gas Producers "Sound and Marine Life Joint Industry Programme", and private donors to the Oregon State University Endowed Marine Mammal Institute.
1555	L.R.P. was funded by the Save Our Seas Foundation and supported by the Manta Trust, the University of Western Australia, and the Australian Institute of Marine Science. Field work was supported by the SOSF-D'Arros Research Centre.
1560 1565	 N.J.P. was funded by Emirates Nature – WWF and multiple supporting agencies. M.P. acknowledges that New Zealand funding was provided by MBIE Endeavour Fund C01X1710 (Ross-RAMP), NIWA SSIF (Coasts & Ocean Centre, programme 4) and NIWA Strategic CAPEX. L.P. and M. Lopez Mendilaharsu were funded by the Convention of Migratory Species, WWF as part of the Trans-Atlantic Leatherback Conservation Initiative, and Peoples Trust for Endangered Species, UK. The Coordenação de Aperfeiçoamento de Pessoal de Nivel Superior (CAPES) provided a grant to M. Lopez Mendilaharsu. N.Q. acknowledges CEECIND/02857/2018; PTDC/MAR/100345/2008 and COMPETE FCOMP-01-0124-FEDER-010580; PTDC/BIA/28855/2017 - COMPETE 1094 POCI-01-0145-
1570	 FEDER-028855. F.R.C. acknowledges the transitory norm contract at the University of Coimbra (DL57/2016/CP1370/CT90) and the projects UIDB/04292/2020, UIDP/04292/2020, granted to MARE, and LA/P/0069/2020, granted to the Associate Laboratory ARNET, financed by the Foundation for Science and Technology (FCT; Portugal).
1575	J.A.R. acknowledges EU INTERREG project FAME (Future of the Atlantic Marine Environment), Fundação para a Ciência e Tecnologia (FCT; SFRH/BPD/63825/2009 and SFRH/BPD/85024/2012), EU project LIFE09 NAT/PT/000041 and by EU INTERREG project FAME 2009-1/089, as well as co-sponsorship by FCT and the European Social Fund (POPH, EU) through post-doctoral grants SFRH/BPD/95372/2013SFRH/BPD/85024/2012 and the strategic program of MARE (UID/MAR/04292/2013 and UID/MAR/04292/2020), and LIFE project "Safe Islands for Seabirds" (LIFE07 NAT/P/000649). A.J. Read acknowledges the U.S. Navy (Atlantic Fleet Forces Command) for funding field work
1580	 and analysis. A.F.R. acknowledges the TOTAL Foundation and TOTAL Muscat Branch. D.R. was supported by Save our Seas Foundation, Project Aware, Royal Geographical Society through (EXERCISE JURASSIC SHARK 2), A1 scuba, downtown aquarium, Azul Marino
1585	 Restaurant, Palapas Ventana, WWF-telcel, PADI, National Geographic, and Cabo Expeditions. R.D.R. acknowledges funding from the Australian Research Council Linkage grant LP140100404 and the Holsworth Wildlife Research Endowment R.R.R. and P.J.N.B. acknowledges funding provided by the National Research Foundation Thuthuka (Grant: 76230) and South African National Antarctic programmes (Grants: 93071 & 110722), through the Department of Science and Technology. Perublic of Science the
1590	Mohamed bin Zayed Species Conservation Fund (project number: 10251290) and the IWC Southern Ocean Research Partnership (IWC-SORP).

F.G.R. acknowledges funding provided by the US Marine Mammal Commission under Grant no. E4047335 and ONR Grant N00014-08-1-1195, the E&P Sound and Marine Life Joint Industry Project of the International Association of Oil and Gas Producers. 1595 D.P.R. acknowledges the Qatar Ministry of Municipality and Environment (QMMOE) and Maersk Oil Research and Technology Centre (MORTC), and was supported by two small grants from the Save Our Seas Foundation. Many thanks to the Save Our Seas Foundation, Al Ghurair Foods and the Emirates Diving Association, Emirates Natural History Group and Le Meridien Al Agah Beach Resort for providing financial support for individual satellite tags. **P.W.R.** acknowledges that northern elephant seal research was supported by the Moore, 1600 Packard, and Sloan Foundations with additional support from the Office of Naval Research and the E&P Sound and Marine Life Joint Industry Project of the International Association of Oil and Gas Producers, Exxon-Mobil and Shell oil. J.P.R. was supported by Juan de la Cierva Formacion program (Ref. FJC2019-040622-I) funded by MCIN/AEI/ 10.13039/501100011033. He also received additional funding from Sustainable 1605 Ocean Alliance for the Artificial Intelligence & Animal Movement (AIAM, ref. D017) project under the SOA Grants program. Funding from Vicenç Mut program of Govern de les Illes Balears. T.L.R. acknowledges the Australian Research Council Linkage Program, LP0989933; Antarctic Science Advisory Committee Program 1144, Sea World Research & Rescue Foundation Inc., 1610 Australian Research Council; and the Scott Foundation. C.A.R. and S.J.P. acknowledge two private trusts, Aqua-Firma, the Shark Foundation, Waterlust, Rufford Small Grant, and the PADI Foundation. Y.R. acknowledges the French Polar Institute Paul Emile Victor (IPEV), the WWF-UK, the PEW Foundation, the Centre National de la Recherche Scientifique (Programme Zone Atelier de 1615 Recherches sur l'Environnement Antarctique et Subantarctique, ZATA), the Agence Nationale pour la Recherche (ANR-2010-BLAN-1728-01), the Fondation Albert II de Monaco, and the Fondation des Treilles. P.M.S. was funded by New Zealand's Foundation for Research, Science & Technology under contracts CO1X0008 and CO1507. 1620 G. Schofield acknowledges Deakin University, Australia; Queen Mary University of London, UK; Swansea University, UK; National Marine Park of Zakynthos, Greece; AXA Research Fund, Boyd Lyon Sea Turtle Fund, British Chelonia Group, People's Trust for Endangered Species, Project Aware, and Thermadap. J.M.S. was funded by the Holsworth Wildlife Research Endowment. 1625 S.A.S. acknowledges TOPP funding (ONR, NSF, Moore, Sloan, and Packard Foundations). **K.S.** acknowledges funding received by the Centre for Ecological Sciences, Indian Institute of Science, Bangalore; the Indian Space Research Organisation/Indian Institute of Science, Bangalore Space Technology Cell; and the International Seafood Sustainability Foundation. 1630 G.L.S. acknowledges major funding from support of the TOPP program of the Census of Marine Life, and was supported by the Alfred P. Sloan Foundation, the Gordon and Betty Moore Foundation, the Packard Foundation, the National Oceanographic Partnership Program of ONR, the United Nations Educational, Scientific and Cultural Organization (UNESCO) World Heritage Program (via the United Nations Foundation and Global Conservation Fund of 1635 Conservation International), the Dr. Earl H. Myers and Ethel M. Myers Oceanographic and Marine Biology Trust, the sponsors of the 2007 Great Turtle Race, Earthwatch Institute and the

National Aeronautics and Space Administration (NASA) through a grant provided by the Applied Sciences Program in the Earth Science Division. M.A. Silva and R.P. acknowledge funds provided by FCT through research grants TRACE-1640 PTDC/MAR/74071/2006, IF/00943/2013/CP1199/CT0001, individual contracts/grants to MAS (FCT-IF/00943/2013) and RP (SFRH/BPD/108007/2015), and the strategic projects MARE (UID/MAR/04292/2019) and Okeanos (UIDB/05634/2020), co-funded by FEDER, COMPETE, OREN, POPH, FSE, and the Portuguese Ministry for Science and Education; the Regional Government of the Azores, FRCT, and the Operational Program AZORES 2020 through research grant MAPCET-M2.1.2/F/012/2011, project M1.1.A/REEQ.CIENTÍFICO 1645 UI&D/2021/010, and contracts to MAS and RP through Fund 01-0145-FEDER-000140 "MarAZ Researchers: Consolidate a body of researchers in Marine Sciences in the Azores" of the European Union. **D.W.S.** further acknowledges additional field research support provided by the Save Our Seas 1650 Foundation (Grant Nos. 45, 87, 308) and the NERC Oceans 2025 Strategic Research Programme. G. Skomal acknowledges the Large Pelagics Research Center (Grant 06-125), Federal Aid in Sport Fish Restoration, NSF (OCE-0825148), the John J. Sacco and Edith L. Sacco Charitable Foundation, the Atlantic White Shark Conservancy, the Massachusetts Environmental Trust, Discovery Communications, National Geographic, and the Woods Hole Oceanographic 1655 Institution. L.L.S. thanks G. Hays for contributing funds to purchase four Argos-GPS tags; the Oceanário de Lisboa for contributing funding to purchase two Argos-GPS tags.. Funding was provided by the FCT SFRH/BD/68717/2010. 1660 J.D. Stewart acknowledges the PADI Foundation (Grant No. 7842), the New England Aquarium, MCAF, Carl F. Bucherer, the Punta Mita Foundation, David Connell, Mary O'Malley, Lupo Dion, CIMEC, and the Gulf of California Marine Program. A. Takahashi acknowledges funding provided by the North Pacific Research Board (contribution number 1612-3), Japan Society for the Promotion of Science KAKENHI Grant Number JP16H02705, and the Arctic Challenge for Sustainability program 1665 (JPMXD130000000) of Japan Ministry of Education, Culture, Sports, Science and Technology. Work was supported by Grant-in-Aid for Scientific Research (20241001 and 24370016). JSPS research grants (19651100 and 19255001), and by a Grant-in-Aid for Scientific Research (Special Promotion) of the Ministry of Education, Culture, Sports, Science and Technology-1670 Japan to Yamashina Institute for Ornithology. P.M.T. acknowledges funding provided by Marine Alliance for Science and Technology for Scotland, Beatrice Offshore Windfarm Ltd, Crown Estate, Highlands & Islands Enterprise, and Moray Firth Offshore Renewables Limited. J. Tomás thanks the European Union Marie Curie FP7 and the Spanish Ministry of Education 1675 and Sciences, and was also supported by project LIFE INTEMARES (LIFE18 NAT/IT/000103) F.V. acknowledges the Foundation for Science and Technology (FCT) for individual grants (CEECIND/03469/2017, CEECIND/03426/2020) and research funds under the project UIDB/05634/2020 and UIDP/05634/2020, and support of the Regional Government of the Azores through the initiative to support Research Centers of the University of the Azores and 1680 through the project M1.1.A/REEQ.CIENTÍFICO UI&D/2021/010.

1685	 M. Vedor was funded by Fundação para a Ciência e Tecnologia (FCT; PTDC/ASP-PES/2503/2020). S.V. thanks the TOPP Program supported by the Sloan, Packard and Moore Foundations, as well as the ONR, the E&P Sound and Marine Life Joint Industry Project of the IAGOP (#JIP 2207-23), UC MEXUS, CONACYT in Mexico and Instituto Politécnico Nacional (IPN) (Project SIP-
	2012006) of Mexico, NSF Office of Polar Programs and Center for Remote Sensing at University of California Santa Cruz for funding and logistic support.
	C.V. was funded by DREAL Bretagne, FEOGA Funds (EU), DREAL Basse Normandie, Région Poitou-Charente, La Compagnie du Vent, and the Parc naturel marin d'Iroise.
1690	S. Wanless was funded by the NERC Centre for Ecology & Hydrology.
	R.S.W. was funded by ONR and Dolphin Quest, Inc.
	S.D.W. acknowledges the Department of Environment and Natural Resources, Conservation Volunteers Australia, Tiwi Land Council, Natural Heritage Trust, Charles Darwin University, and the Australian Government.
1695	B.W. acknowledges the Australian Antarctic Division.
	N.E.W. acknowledges that tags were part of WWF-Australia's Flatback Whereabouts Project funded by WWF, Factorie, and Winnifred Violet Scott Trust, supported by the Gudjuda Aboriginal Reference Group.
1700	 D.N.W. acknowledges the Volgenau Foundation, Mudge Foundation, BOEM, Stellwagen Bank National Marine Sanctuary, and the National Marine Sanctuary Foundation. F.C.W. was funded by the UK Natural Environment Research Council (NERC) through a University of Southampton INSPIRE DTP Studentship. L.J.W. acknowledges funding by the UK Department for Energy and Climate Change (DECC).
1705	 J.C.X. acknowledges the strategic program of MARE (Marine and Environmental Sciences Centre), financed by the Foundation for Science and Technology (UIDB/04292/2020), through the grants Investigator FCT program (IF/00616/2013), PTDC/BIA-BDE/64539/2006 and SFRH/BPD/28879/2006. T.Y. was partially supported by the Japan Society for the Promotion of Science research grants (19651100, 19255001) and Grant-in-Aid for Scientific Research (Special Promotion) of the
1710	Ministry of Education, Culture, Sports, Science and Technology-Japan.
	D.J.Y. acknowledges funding provided by Polar Continental Shelf Program, Ocean Tracking Network, Fisheries and Oceans Canada, University of Windsor, Ontario Graduate Scholarships, W. Garfield Weston Foundation and Natural Science and Engineering Research Council of Canada.
1715	P.M.Z. acknowledges the United Nations Office for Project Services, GEF Humboldt for providing tags and funding for the expedition
1720	Author contributions: All authors contributed to aspects of fieldwork, animal tagging, data collection, data formatting, and/or contribution of tools (full details provided in Supplementary Text: "Supplementary Author Contributions")
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	Investigation: All authors
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	Writing – review & editing: All authors
1730	Competing interests: Authors declare that they have no competing interests.
	Data and materials availability: All data, code, and materials needed to reproduce this analysis are available at <u>https://doi.org/10.5061/dryad.x95x69ptv (83)</u> . Data obtained from literature review are presented in supplementary materials. Sources of environmental data collated from online databases are described in supplementary materials.
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1745 References (112-135)

Fig. 1. Tracked movements of marine megafauna at the global scale.

A) Map of the total number of 12,794 unique individual track locations in the global dataset at 1° resolution showing the global coverage of 71.7% of the global ocean. B) Maps per taxon
1750 showing the number of unique individual track locations within each 1° grid-cell. From top left to bottom right, maps per taxon show 6324 individual tracks for 39 species of flying birds, 749 for cetaceans including 11 whales and 3 delphinid species, 1760 for fishes including 23 shark species, 2 manta rays, and 1 ocean sunfish, 1324 for 6 species of penguins, 65 for polar bears, 1698 for 16 species of seals, 28 for sirenians including dugongs and West Indian manatees, and 846 for all 7 sea turtles. The latitudinal and longitudinal coverage of tracked data is displayed in fig. S1. For reference, the first position obtained for each tracked individual (i.e., representing tagging locations), as well as captured and expected global biodiversity are given in fig. S2. Maps showing the spatial extent of space use per species at 1° resolution can be seen in the data repository.

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Fig. 2. Global space-use of marine megafauna and time spent in different behaviours.

Fractions of area (A) and time (B) used by animals globally (left plots), within and outside exclusive economic zones (EEZs) (middle plots), and within and outside existing marine protected areas (MPAs) (right plots), showing how much of the movements corresponded to
detected migratory corridors or residency. Results are shown across all species together (top bar) and for each taxon (as displayed in legend). For each taxon, the light grey portion in the bars indicates movement where no behaviours were detected. Species in each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (*Ursus maritimus*), seals, sirenians (i.e., dugongs and manatees), and turtles. C) Map of detected migratory corridors, residence areas and both corridors and residencies across taxa. Grey indicates grid-cells where tracking data were available but no specific behaviour was identified for any taxon. Light blue areas depict regions where we did not have tracking data. Maps of detected behaviours per taxon can be seen in fig. S9.

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Fig. 3. Increase in area protection to 30% will leave ~60% of IMMegAs exposed to major anthropogenic threats.

A) Maps depicting average threat intensities for major anthropogenic threats with a global footprint: (from top to bottom) fishing, shipping, plastic pollution and sea surface temperature (SST) warming. Displayed with an orange colour palette are the threat intensities occurring 1780 inside IMMegAs, while a grey colour palette is used to show the threat intensities outside IMMegAs. Note that we considered noise to be ubiquitous, as no noise dataset is currently available at the resolution needed for a global analyses. B) Maps showing how much the increase in marine protected areas (MPAs) from the current 8% (purple) to 30% (green) would cover 1785 from our prioritization of IMMegAs detected from movement data (top map) and from our model predictions (bottom results). Note that coverage by MPAs only translates into protection from the anthropogenic threats considered if they are designated with the highest level of protection (i.e., with no activities allowed), and even then MPAs could only be effective for protection from fishing and shipping, leaving plastic and warming threats to continue to affect 1790 species. In addition to the increase in the current extent of MPAs, the introduction of mitigation strategies will assist in reducing the impact of existing threats and therefore the likelihood of human-induced extinctions.

- 1794 Table 1. Evidence of impacts from overlap of marine megafauna with anthropogenic
- 1795 threats. Examples of the range of impacts derived from the overlap of marine megafauna
- 1796 with anthropogenic threats such as climate warming, plastic pollution, shipping, noise
- 1797 pollution, and fishing. SST: sea surface temperature; UV: ultraviolet.

	BIRDS (FLYING)	CETACEANS	FISHES	PENGUINS	POLAR BEAR	SEALS	SIRENIANS	TURTLES
	Decreased survival	UV damage	Habitat shift	Reduced prey	Habitat contraction	Habitat shift	Reduced food	Sex bias
CLIMATE	Impacted survival & population growth rate of black-browed albatross juveniles with SST changes (84)	Increased skin lesions on whale related with increased UV irradiance (85)	Reduced counts of Scalloped hammerhead sharks <i>Sphyrna lewini</i> associated with rise in SST (<i>86</i>)	Decreased population size for penguin prey species with climate change (87)	Contraction of polar bear's habitat in the Arctic linked to long-term sea ice loss (88)	Decreased survival of southern elephant seal due to effects of sea ice dynamics on access to foraging (89)	Reduced dugong density by ~70% due to seagrass die- off triggered by an extreme heat wave (90)	Female-biased turtle populations linked to warming temperatures (91)
	Ingestion	Ingestion	Ingestion	Ingestion		Entanglement	Ingestion	Ingestion
PLASTIC	Death of shearwater and northern gannet due to plastic ingestion (92)	Stranded sperm whale stomachs with large amounts of plastic debris (93)	Threatened filter- feeding elasmobranchs by microplastic (94)	Plastic ingestion may have caused death (95)	-	Mortality of fur seals due to entanglement in marine debris (96)	Death of West Indian manatees from ingestion of plastic debris (97)	50% probability of mortality when turtles ingest pieces of plastic (98)
7 8	Habitat loss	Ship strike	Ship strike	Noise effects	Ship strike	Propeller strike	Ship strike	Ship strike
SHIPPING	Habitat loss for Common Eider's avoiding shipping traffic (99)	Increased ship strikes with humpback whales in shipping lanes (39)	Mortality of whale sharks correlated with risk of collision with ships (41)	Population collapse concomitantly with increase in noise (100)	Increased vulnerability of polar bears to vessel strike (101)	Propeller strikes affect harbor seals (102)	Death of manatees due to boat collisions (103)	Decreased survival of green turtles due to boat strikes (104)
		Behav. change			Disturbance	Physical damage	Behav. change	
NOISE	-	Change in humpback whales foraging activity due to ship noise (105)	-	-	Disturbance of maternal dens due to seismic surveys (106)	Temporary hearing loss of grey and harbor seals around the British Isles (107)	Reduced foraging habitat for manatees due to boat noise (108)	-
	By-catch	By-catch	Mortality	Reduced prey		Entanglement	Entanglement	By-catch
FISHING	High bycatch of seabirds in longline fisheries (38)	Higher rates of dolphin bycatch in a trawl fishery (109)	Greater mortality of pelagic sharks where sharks have higher exposure to longline fisheries (62)	Decreased population size of prey species with increased fishing of Antarctic Krill (<i>87</i>)	-	Increased entanglement of Cape fur seals associated with fishing (110)	Manatee mortalities from entanglement in fishing gear (111)	High levels of turtle bycatch in fishing gear hotspots (<i>37</i>)

1799 Table 2. Summary of the logistic modelling inputs and results per taxon

Results of the generalized linear models relating the probability of a grid-cell to be used as residence or for migratory behaviours with the set of environmental variables included in each model. Shown are the results for the highest ranked model according to the weight of the Akaike's Information Criteria (*wAIC*), as well as the number of parameters (k), the percentage of deviance explained (pcdev) and Kappa. Grey indicates the models not used to estimate the important marine megafauna areas (IMMegAs) derived from our modelling predictions (as presented in Fig. 3 and fig. S11). Species in each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (*Ursus maritimus*), seals, sirenians (i.e., dugongs and manatees), and turtles.

Input					Results									
Taxon	Numb	er of grid-cell	s with:]	Resid	lence Be	ehaviou	•	I	Migra	atory Be	haviour	•	
	Presence	Residency	Migration	Model	k	wAIC	pcdev	Kappa	Model	k	wAIC	pcdev	Kappa	
Birds	35,875	13,448	9,128	2	19	1.000	4.13	0.22	2	19	1.000	11.19	0.33	
Cetaceans	4,397	1,501	1,758	2	19	1.000	16.52	0.44	2	19	0.980	12.62	0.29	
Fishes	15,648	4,346	4,252	2	19	1.000	14.44	0.38	2	19	1.000	12.56	0.30	
Penguins	1,385	446	452	1	17	1.000	13.62	0.4	2	19	1.000	40.16	0.56	
Polar	1,124	451	803	2	14	0.005	24 78	0.22	2	14	1 000	27 78	0.48	
bear				Δ	14	0.995	24.70	0.55	L	14	1.000	27.78	0.40	
Seals	11,358	5,510	7,175	2	19	1.000	3.12	0.22	2	19	1.000	14.91	0.30	
Sirenians	114	27	0	-	-	-	-	-	-	-	-	-	-	
Turtles	10,360	3,462	3,370	3	7	1.000	7.71	0.28	2	19	1.000	5.18	0.17	

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1817	Ana M. M. Sequeira ^{1,28} , Jorge P. Rodríguez ^{3,4,5,68} , Sarah A. Marley ⁷ , Hannah J. Calich ^{1,8} , Mirjam
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1828	Michael L. Berumen ⁶³ , Sandra Bessudo ⁶⁴ , Natalia P. A. Bezerra ^{65,66} , Antonin V. Blaison ^{67,68} ,
1829	Gabriela S. Blanco ⁶⁹ , Barbara A. Block ⁷⁰ , Mark Bolton ⁷¹ , Mark E. Bond ⁷² , Ramón Bonfil ^{73,74,73} ,
1830	Camrin D. Braun ⁷⁰ , Annette C. Broderick ⁷⁷ , Michael de L. Brooke ⁷⁰ , Annabelle M. L. Brooks ⁷⁷ ,
1031	Campana ⁸³ Hamish A. Camphell ⁸⁴ Richard A. Camphell ⁸⁵ Aaron Carlisle ⁸⁶ Ruth H
1832	Carmichael ^{87,88} Gemma Carroll ⁸⁹ Paolo Casale ⁹⁰ Filine R Ceia ⁹¹ Demian D Chapman ^{92,93}
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1836	Eric E. G. Clua ^{104,105,106} , Jesse E. M. Cochran ⁶³ , Rochelle Constantine ¹⁰⁷ , Robert W. Cooper ¹⁰⁸ ,
1837	Estelle Crochelet ⁶⁷ , Michelle Cronin ^{58†} , Eduardo Cuevas ^{109,110,111,112} , Kayla P. DaCosta ^{87,88} ,
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1846	Sabrina Fossette ^{10,151} , Malcolm P. Francis ¹⁴⁵ , Ari S. Friedlaender ¹⁰⁰ , Miguel Furtado ¹³ , Austin J.
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1914	
1915	
1916	The PDF file includes:
1917	
1918	Materials and Methods:
1919	 Fieldwork and deployment of tracking devices
1920	- Animal ethics information
1921	- Tracking data collection and processing
1922	 Addressing tracking data biases
1923	- Detection of key movement behaviours
1924	- Statistical modelling
1925	- Optimisation algorithm
1926	Supplementary Acknowledgements
1927	Supplementary Author Contributions
1928	Figs. S1 to S17
1929	References (112-135)
1930	

1931 Materials and Methods

1932 Fieldwork and deployment of tracking devices

1933 1934 Birds

1935 Birds were all caught at nest sites either whilst incubating or attending chicks, except for some 1936 northern gannets (Morus bassanus, immature birds at the main colony), Trindade petrels 1937 (Pterodroma arminjoniana, non-breeding adult birds resting at the main colony), and great 1938 shearwaters (Ardenna gravis, attracted to a vessel at-sea using bait). Birds were captured using 1939 noose poles, crook poles, drop traps, net launchers, nets (landing, mist, purse or handheld), or 1940 removed by hand from their burrows. Tags were typically attached to the auxiliary leg band or 1941 taped to the mantle, scapular, dorsal contour, or tail feathers. Chest or leg-loop harnesses were 1942 used for herring gulls (Larus argentatus), ivory gulls (Pagophila eburnea), some Ross's gulls 1943 (Rhodostethia rosea), and some northern fulmars (Fulmarus glacialis). For great shearwaters, 1944 tags were attached dorsally using four subcutaneous Prolene sutures. In all cases, total instrument 1945 mass was <5% of body mass to minimise effects on flight efficiency and all birds were handled

- 1946 for less than 20 minutes.
- 1947

1948 Cetaceans

1949 Smaller cetaceans (e.g., beluga - Delphinapterus leucas, bottlenose dolphins - Tursiops

1950 *truncatus*, narwhal - *Monodon monoceros*) were captured using seine or stationary nets. The

animals were then brought to the surface, disentangled, and secured using hoop nets and loop

ropes. In the case of bottlenose dolphins, animals were brought aboard the research vessel as part

1953 of capture-release health assessments. Tags were attached using nylon pins attached to the dorsal 1954 ridge or fin. Killer whales (*Orcinus orca*) were targeted from shore using crossbows and tags

1954 Huge of fin. Kiner whates (*Orethus orea*) were targeted from shore using crossbows a 1955 were attached to the dorsal fin using subdermal darts. Other cetaceans, such as blue -

1956 Balaenoptera musculus, bowhead - Balaena mysticetus, gray - Eschrichtius robustus, humpback

1957 - Megaptera novaeangliae, pilot - Globicephala macrorhynchus and G. melas, right - Eubalaena

1958 glacialis and E. australis, and sei whales - Balaenoptera borealis, were approached using a small

research vessel. Tags were deployed using crossbows, air-powered applicator systems, or long

1960 fibreglass poles. Tags were attached to the dorsal fin (either anterior-to or at-the-base-of) using

subdermal anchors or barbs and petals, which were sterilised and/or treated with antibiotic

1962 coatings prior to deployment.

- 1963
- 1964 Fishes

Fish, mostly sharks, were typically captured with baited hooks, bagan lift nets, or in purse-seinenets, then brought alongside the vessel and restrained in a sling or with straps, secured to a

raisable platform, or taken aboard for tagging. If brought aboard, fish were on deck an average of

approximately 3 minutes; the exceptions to this were white sharks - *Carcharodon carcharias*

- (average duration of restraint: 12 mins) and tiger sharks *Galeocerdo cuvier* (some were placed
 in tanks with running seawater and moved to deeper isobaths as part of a shark attack mitigation
- 1970 In tanks with fulling seawater and moved to deeper isobatils as part of a shark attack initigatility strategy). Manta rays *Mobula birostris* and *M. alfredi*, and some copper *Carcharhinus*
- *brachyurus*, Galapagos *Carcharhinus galapagensis*, scalloped hammerhead *Sphyrna lewini*,
- 1973 whale *Rhincodon typus*, and white sharks were tagged whilst free-swimming in the water

1974 1975 1976 1977 1978 1979 1980 1981 1982 1983	column using pneumatic spear guns or rubber-propelled hand spears. The majority of sunfish (<i>Mola mola</i>) and some porbeagle sharks (<i>Lamna nasus</i>) were captured as bycatch in fisheries targeting tuna. Tags were typically attached using a tether affixed to a dart, which was implanted in the dorsal musculature or anchored to the first (or second in bluefin tuna - <i>Thunnus thynnus</i> ; removed from analyses) dorsal fin. For some sunfish, tags were attached to the base of the caudal fin. For some of the blue - <i>Prionace glauca</i> , bull - <i>Carcharhinus leucas</i> , mako - <i>Isurus oxyrinchus</i> and <i>I. paucus</i> , sandbar - <i>Carcharhinus plumbeus</i> , scalloped hammerhead, silky - <i>Carcharhinus falciformis</i> , tiger, whale, and white sharks, tags were attached to the first dorsal fin using metal bolts, neoprene and high-carbon steel washers, and steel nuts. For some white sharks, tags were mounted on a custom-built spring clamp that was placed on the first dorsal fin.
1984 1985	Penguins
1986 1987 1988	Penguins were captured and released on land at nesting sites. Tags were attached to the dorsal plumage using waterproof tape and/or epoxy glue, and in some cases secured under a bed of feathers using a small cable tie.
1989 1990	Polar bears
1991 1992	Adult female polar bears (<i>Ursus maritimus</i>) were located via helicopter and immobilised with a rapid-injection dart. Tags were attached using satellite collars.
1993 1994	Seals
1995 1996 1997 1998	Seals were approached whilst onshore or in shallow waters surrounding haul-out sites and captured using hoop nets, tangle nets, beach seine nets, and/or remote syringe darts. Once captured, seals were manually restrained, sedated, or anaesthetised. Tags were attached to the head or along the dorsal midline using quick-setting epoxy glue.
1999 2000	Sirenians
2001 2002 2003 2004 2005 2006	Manatees (<i>Trichechus manatus</i>) were located via an aerial observer and individuals were captured in a net deployed from a specialised capture boat. Dugongs (<i>Dugong dugon</i>) were captured using a 'rodeo' technique, where a personal watercraft is used to closely pursue an individual dugong until fatigued. The dugong is then caught around the peduncle region by a catcher leaping off the boat, and the animal is restrained at the water surface by several people. For all sirenians, tags were tethered to the animal using a peduncle belt.
2007 2008	Turtles
2009 2010 2011 2012 2013 2014	Turtles were primarily adult females captured at nesting beaches after a successful nesting event. In some cases, adult and juvenile turtles were captured at sea (both in the vicinity of nesting beaches or at foraging grounds) using tangle nets, dip nets, a "rodeo" technique, or by hand as they rested at the surface. Some turtles were found stranded or were incidentally captured by local fishers, then handed into conservation organisations for tagging and release. For hard- shelled turtles, tags were attached to the carapace or head with quick-setting epoxy glue, a

2015 fiberglass and polyester resin, or in the case of flatback turtles (*Natator depressus*), by using a

- 2016 specially-designed harness. Leatherback turtles (*Dermochelys coriacea*) were tagged via direct
- 2017 attachment surgical technique (tags were directly attached by drilling into the central-dorsal ridge
- 2018 and affixing with nylon or metal ties), tow technique (hole drilled in caudal peduncle and tag
- 2019 towed), or harness technique. Where post-hatchlings were used, they were collected from the
- 2020 nest, reared by head-starting programs, and then selected for tagging based on their size and
- 2021 swimming abilities. Post-hatchlings were tagged using an acrylic-silicone-neoprene attachment
- 2022 method, which for larger individuals sometimes also included drilling through the keratin part of
- 2023 the carapace crest and securing the tag with nylon ties.

2024 Animal ethics information

2025 Data providers obtained all licenses and ethical permissions required for data collection in their 2026 jurisdictions and ensured that each animal was handled and tagged by trained personnel. Details 2027 per taxon are presented below with name initials indicating the responsible co-author.

- 2028
- 2029 Birds (flying)
- Tagging of black-browed albatrosses (*Thalassarche melanophris*) at Diego Ramirez Islands was
 conducted under a permit provided by the Chilean Antarctic Institute (J.A.A.).
- 2032 Audouin's gull (Ichthyaetus audouinii) tagging was conducted with permission from the Catalan
- 2033 and Balearic Islands Governments, and Scopoli's shearwater (*Calonectris diomedea*) tagging 2034 was conducted with permission from the Balearic and Valencian Governments, as well as the
- 2035 Spanish Government (**J.M.A.**).
- The sooty tern (*Onychoprion fuscatus*) tracking project in Seychelles was approved by the Seychelles Bureau of Standards and supported by the owners of Bird Island (**C.F.**).
- 2038 Tagging procedures on little penguins (Eudyptula minor), crested terns (Thalasseus bergii), and
- 2039 short-tailed shearwaters (Ardenna tenuirostris) off South Australia were conducted under
- 2040 approval by the South Australian Department of Primary Industry and Regions (PIRSA) Animal
- 2041 Ethics Committee (32-12) and the South Australian Department for Environment and Water (DEW) (S. E. C.)
- 2042 (DEW) (Scientific Permit A24684) (**S.D.G.**).
- 2043 Broad-billed prion (*Pachyptila vittata*) fieldwork on Rangatira was conducted with the
- 2044 permission and cooperation of the New Zealand Department for Conservation and would not
- 2045 have been possible without the support of the Chatham Island Area Office. Northern gannets
- 2046 were ringed and loggers deployed with permits and approval from the British Trust for
- 2047 Ornithology (BTO) and Scottish Natural Heritage (W.J.G.).
- 2048 All tracking of northern gannets, razorbills (Alca torda), Atlantic puffins (Fratercula arctica),
- 2049 and Manx shearwaters (*Puffinus puffinus*) in the Republic of Ireland were approved by the
- 2050 University College Cork (UCC) Animal Ethics Committee (2013/032 and 2019/001) and
- 2051 conducted under permits by the BTO (C/6143) and Irish National Parks and Wildlife Service
- 2052 (26/2010, 011/2013, 018/2014, 016/2015, 025/2016, 082/2017, C051/2011, C116/2012, 2052 (2012) (
- 2053 C039/2013, C075/2014, C087/2015, C100/2016, C87/2017) (**M.J.**).
- 2054 Barau's petrel (*Pterodroma baraui*) tracking work was authorized by Centre de Recherches sur
- 2055 la Biologie des Populations d'Oiseaux (CRBPO) permit number PP609, Ethic Committee of
- 2056 Réunion Island, Parc National de La Réunion, and direction de l'environnement, de
- 2057 l'aménagement et du logement de La Réunion (DEAL-Réunion). Red-tailed tropic bird
- 2058 (*Phaethon rubricauda*) tagging was authorized by Permit Le Corre PP616, Terres Australes et
- 2059 Antarctiques Françaises (TAAF), Mauritius National Park, and Madagascar National Parks.
- 2060 White-tailed tropicbird (*Phaethon lepturus*) tracking was conducted with research approval by 2061 CRBPO (PP616) and the Seychelles Bureau of Standard (SBS). Sooty tern tagging was
- 2061 authorized by PP616 M. Le Corre, Seychelles Bureau of Standard, and TAAF. Wedge-tailed
- 2062 shearwater (*Ardenna pacifica*) tagging was authorized by CRBPO permit PP616, Ethical
- 2064 committee of Réunion Island, Institutional Authorizations from DEAL-Réunion, Conservatoire
- 2065 du Littoral Réunion, Mauritius National Parks and Conservation Service, and Seychelles Bureau
- 2066 of Standard (**M.L.C.**).

2067 Common eider (Somateria mollissima) tagging was conducted under Environment Canada 2068 (ECCC) Animal Care Permits, Canadian Wildlife Service (CWS) Scientific Permit NUN-SCI-2069 04-02, and Nunavut Wildlife Research Permit WL1028. Herring gull tagging was conducted 2070 under Nunavut Wildlife Research Permit WL2008-1028; CWS Scientific Permit NUN-SCI-08-2071 04, SC2761; and ECCC Animal Care permits EC-PN-08-026. Ivory gull tagging was conducted under CWS Banding Permit number 10694; CWS Scientific Permit NUN-SCI-09-02; and 2072 2073 Nunavut Wildlife Research License WL2010-032. Northern fulmar collections were in 2074 accordance with Canadian Council on Animal Care guidelines, and were conducted under the 2075 following permits: research (NUN-SCI-03-02, WL000190, WL000714), animal care 2076 (2003PNR017, 2004PNR021, 2005PNR021), and land use (59A/7-2-2). Parasitic jaeger 2077 (Stercorarius parasiticus) tagging was conducted under CWS Banding Permit 10694; Animal 2078 Care EC-PNR-11-020, Scientific Permit NUN-SCI-09-01, and Territorial Permit WL 2010-042. 2079 Ross's gull tagging was conducted under CWS Banding Permit 10694; Animal Care Permit EC-2080 PNR-11-020; Scientific Permit NUN-SCI-09-01; and Territorial Permit WL 2010-042. Sabine's 2081 gull (Xema sabini) tagging was conducted under permits CWS Animal Care EC-PN-11-020, 2082 CWS Scientific Permit NUN-SCI-09-01, Government of Nunavut Wildlife Research Licence 2083 WL 2010-042, Nunavut Water Board licence 3BC-TER0811, Indian and Northern Affairs Land 2084 Use Reserve 068H16001, and CWS Banding Permit 10694. Thick-billed murre (Uria lomvia) 2085 tagging was conducted under Canadian scientific and access permits (NUN-SCI-08-55, NUN-2086 MBS-12-03, NUN-SCI-12-04, WRP2013040), banding permit (10694, 10322), and animal care 2087 (0800AG01) (M.L. Mallory).

Tagging work followed the ethical standards set out by the Mauritian Wildlife Foundation and its
partner and consulting organisations, the North of England Zoological Society, the Durrell
Wildlife Conservation Trust, and the International Zoo Vet Group (M.A.C.N.).

2091 Permission to capture and tag Ascension frigatebirds (*Fregata aquila*) was granted by the

2092 Conservation Department of the Ascension Island Government. The attachment of devices met

2093 the ethical guidelines of the Special Methods Panel of the BTO. King eiders (Somateria

2094 *spectabilis*) were handled with approval by the University of Alaska Fairbanks Institutional

2095 Animal Care and Use Committee (IACUC) (protocol #05-29) and CWS Animal Care Committee

2096 (permit #PNR007). Masked booby (*Sula dactylatra*) tagging was carried out under permission

2097 and with collaboration of the St Helena Environmental Management Directorate. The capture 2098 and handling of birds and attachment of unconventional marks was carried out under licence

and handling of birds and attachment of unconventional marks was carried out under licence
 from the BTO. Permission to capture and tag birds was granted by the Environmental

2100 Management Directorate on St Helena. The attachment of GPS devices met the ethical guidelines

2101 of the Special Methods Panel of the BTO. Tagging of Murphy's petrels (*Pterodroma ultima*)

2102 followed all applicable international, national, and/or institutional guidelines for the care and use

2102 followed an applicable international, national, and/or institutional guidelines for the care and use 2103 of animals. Permission to access Henderson Island in order to conduct scientific research in 2015

2104 was granted by the Government of the Pitcairn Islands (**S. Oppel**).

2105 Tagging work was conducted under approval by the Portuguese Government Instituto de

2106 Conservação da Natureza e Florestas (ICNF) under licenses 188/2010/ CAPT, 152/2011/CAPT,

2107 101/2012/CAPT, 99/2013/CAPT, 203/2014/CAPT, 169/2015/CAPT, and 89/2011/CAPT

2108 (V.H.P.).

2109 Tagging work was authorized by the Government of South Georgia and the South Sandwich

2110 Islands (**R.A.P.**).

- 2111 Tagging procedures were conducted under approval by the Portuguese Government ICNF
- 2112 (permit 89/2011/CAPT) and in compliance with Portuguese laws No. 140/99, No. 49/2005, No.
- 2113 316/89, and No. 180/2008 (J.A.R.).
- 2114 Buller's albatross (Thalassarche bulleri) tagging was approved by the Southland Conservancy,
- 2115 Department of Conservation, New Zealand (**P.M.S.**).
- 2116 Sooty shearwater (Ardenna grisea) ethics was approved by the IACUC at the University of
- 2117 California Santa Cruz and approval for the research was provided by the Whenua Hou
- 2118 Management Committee, Rakiura Titi Islands Administering Body, and Southland Department
- 2119 of Conservation in New Zealand. Black-footed (*Phoebastria nigripes*) and Laysan albatross
- 2120 (Phoebastria immutabilis) tagging in the Hawaiian Islands was approved by the University of
- 2121 California Santa Cruz and San Jose State University IACUCs under Master Bird Banding permit
- 2122 23411. Laysan albatross tagging on Guadalupe Island, Mexico was approved by University of
- 2123 California Santa Cruz IACUC under Master Banding Permit 20768.Western gull (Larus
- 2124 occidentalis) tagging was conducted under permission granted by Año Nuevo State Park,
- 2125 California State Parks, California Department of Fish and Wildlife, and the US Fish and Wildlife
- 2126 Farallon Islands National Wildlife Refuge (SUP# 81641). All research protocols were approved
- 2127 by the San Jose State University IACUC (protocol 979) (S.A.S.).
- 2128 Common murre (*Uria aalge*) field work was conducted under Kukulget Inc. land crossing
- 2129 permits, University of Alaska Fairbanks IACUC protocol #471022, US Fish and Wildlife Service
- 2130 (USFWS) scientific collection permit #MB70337A, A. Kitaysky's Master Banding permit
- 2131 #23350, and Alaska Department of Fish and Game's permits #19-140, 18-131, 17-104, 16-089.
- 2132 Streaked shearwater (*Calonectris leucomelas*) tagging procedures were approved by the Animal
- 2133 Experimental Committee of the University of Tokyo and conducted in accordance with the
- 2134 Guidelines for the Care of Experimental Animals, with fieldwork conducted under permits from
- 2135 the Ministry of the Environment and the Agency for Cultural Affairs. Thick-billed murre tagging
- 2136 was conducted under Kukulget Inc. land crossing permits, UAF IACUC protocol #471022,
- 2137 USFWS scientific collection permit #MB70337A, A. Kitaysky's Master Banding permit #23350,
- 2138 and Alaska Department of Fish and Game's permits #19-140, 18-131, 17-104, 16-089. (A.
- 2139 Takahashi).
- 2140 Capture and tagging of Northern fulmar in Scotland was carried out under licences from the
- 2141 BTO (Licence No: AO/4939) and UK Home Office (Licence No: PIL 60/698) following review
- 2142 by the University of Aberdeen ethics committee (**P.M.T.**).
- 2143 Northern gannet capture and tagging on St Kilda was carried out with permission from the
- 2144 National Trust for Scotland and Scottish Natural Heritage and under licence from BTO (Licence
- 2145 No: A2332 with a specific unconventional methods endorsement) (S. Wanless).
- 2146 Tagging procedures were conducted with approval from the US Department of the Interior
- 2147 #21963, Massachusetts Division of Fisheries and Wildlife #058.19SCB, Stellwagen Bank
- 2148 National Marine Sanctuary Permit # SBNMS-2019-001, and the Long Island University IACUC
- 2149 (**D.N.W.**).
- 2150 Northern gannet capture and tagging was carried out under licences from the BTO and Natural
- 2151 England, with approval of the Royal Society of the Protection of Birds (L.J.W.).
- 2152 Tagging work was conducted with permits from the Ministry of the Environment: No.060609001
- 2153 for Sangan Island and No.18–340 for Mikura Island (T.Y.).

- 2154
- 2155 Cetaceans
- 2156 Tagging was undertaken under US National Marine Fisheries Service (NMFS) Scientific
- 2157 Research Permits No. 17096, 731-1774, and 15330. Tagging was undertaken under protocols
- 2158 approved by the Cascadia Research Collective IACUC (**R.W.B.**).
- 2159 Tagging was conducted under University of Auckland Animal Ethics AEC001587, New Zealand
- 2160 Department of Conservation Permit #44388-MAR, and approval from local Maori tribes (iwi)
- 2161 Ngāti Kuri and Te Aupōuri (**R.C.**).
- 2162 Tagging was undertaken with the permission of the Environment Department of the province
- 2163 Sud of New Caledonia and of the Government of New Caledonia under permits 383-
- 2164 2010/ARR/DENV, 33313-2010/ARR/DENV, 3616-2011/ARR/DENV, 3157-2012/ARR/DENV,
- 2165 1045-2014/ARR/DENV, 151-2015/ARR/DENV, 1105-2016/ARR/DENV, 899-
- 2166 2017/ARR/DENV, 2220-2018/ARR/DENV, 2016-1391/GNC, 2017-1107/GNC and 2018-
- 2167 923/GNC (**C. Garrigue**).
- 2168 Tagging procedures were approved by Fisheries and Oceans Canada (DFO) Freshwater Institute
- 2169 Animal Care Committee (AUP # FWI-ACC-2002, 2003, 2004, 2005, 2006 and 2007) and under
- 2170 DFO License to Fish for Scientific Purposes #S-02/03 to 05/06-1019-NU and #S-12/13-1024-
- 2171 NU, S-13/14-1009-NU and S-16/17 1005-NU (S.H.F.).
- 2172 Tagging was conducted under permits 11-101/VP/MPEEIA:SG and 12-100/VP/MPEEIA:SG
- 2173 issued by the Secretary-General of the Union of the Comoros, permits 105/DEAL//SEPR/2012
- and 148/DEAL/SEPR/2012 issued by Direction de l'Environnement, de l'Aménagement et du
- 2175 Logement de Mayotte, and permit FR1397600001-E issued by Direction de l'environnement, de
- 2176 l'aménagement et du logement (DEAL) Mayotte (S.F.).
- 2177 Tagging was conducted under NMFS permits (numbers 14907, 14809, and 14856) and ACA
- 2178 Permits (2009-013 and 2015-011). All animal work was approved and conducted under Duke
- 2179 University IACUC A049-122-02 and the Oregon State University Animal Care and Use Protocol
- 2180 (ACUP) 4513 (**A.S.F.**).
- 2181 Beluga tagging was carried out with Animal Care Approval and Research Permits issued by the 2182 Canadian Government (**M.O.H.**).
- 2183 Deployment of satellite tags on southern right whales at the Head of Bight, South Australia were
- 2184 conducted under approval by the South Australian Department of Primary Industries and
- 2185 Regions (PIRSA) Animal Ethics Committee (32-12), and under the following permits: PIRSA
- 2186 Fisheries Exemption (ME9902712), Department of Environment Water and Natural Resources
- 2187 (DEWNR) Permit and Licence to Undertake Scientific Research (A24684-12), Environment
- 2188 Protection and Biodiversity Conservation Act Cetacean Permit (20014-0004), Access to
- 2189 Biological Resources in a Commonwealth Area for Non-commercial Purposes (AU-COM2014-
- 2190 248), Approval for Activity in Commonwealth Marine Reserve (CMR-14-000196) and DEW
- 2191 Marine Parks Permit (MO00024-2) (A.I.M., S.D.G., and R.H.).
- 2192 Beluga tagging was carried out under Animal Use Protocol permit number FWI-ACC-2015-018
- and DFO license S-12/13-1022-NU. Narwhal tagging was carried out under Animal Use
- 2194 Protocol number FWI-ACC-2016-030 from the DFO Animal Care Committee (under the
- 2195 Canadian Council on Animal Care) and a DFO License to Fish for Scientific Purpose License S-
- 2196 16/17-1037-NU (**M. Marcoux**).

- 2197 Sei whale fieldwork and tagging was approved by the Regional Directorate of the Environment/
- 2198 Regional Government of the Azores under research permit 7/CN/2005, issued to the Department
- 2199 of Oceanography and Fisheries of the University of the Azores (E.O.).
- 2200 Whale tagging was authorized by the NMFS under permit numbers 841 (for blue, bowhead,
- 2201 gray, and humpback whales), 369-1440 (for blue, fin *Balaenoptera physalus*, gray, humpback
- and northern right whales), and 369-1757 (for blue, gray, and southern right whales). Tagging in
- 2203 Mexican waters was conducted under permits issued by the Secretaría de Medio Ambiente y 2204 Recursos Naturales, Mexico (permit number DOO 02.8319 and SGPA/DGVS 0576). Southern
- right whale tagging was also authorised under a permit issued by the South African Department
- 2206 of Environmental Affairs and Tourism in terms of Regulation 58 of the Marine Living Resources
- 2207 Act (no. 18 of 1998). Sperm whale (*Physeter macrocephalus*) tagging was conducted under
- 2208 permits # 08159 and SGPA/DGVS 01102 by the Secretaría de Medio Ambiente y Recursos
- 2209 Naturales of Mexico, and the NMFS under permit numbers 369-1757. For all eight species,
- 2210 research was approved by the Oregon State University IACUC (**D.M. Palacios** and **B.M.**).
- Tagging was approved by the University of Pretoria's Ethics Committee (EC023-10; EC077-15)
- and permitted by the Prince Edward Islands Management Committee (PEIMC 17/12, 1/2013 and
- 2213 1/2014) (**R.R.R.** and **P.J.N.B.**).
- 2214 Blue, fin and sei whale fieldwork and tagging were approved by the Regional Directorate of the
- 2215 Environment/Regional Government of the Azores, under research permits: 20/2009/DRA (blue,
- fin and sei whales), 16/2010/DRA (blue and fin whales), 51/2011/DRA (blue and fin whales),
- 2217 30/2015/DRA (blue whale), 37/2016/DRA (blue whale), 31/2012/DRA (fin whale),
- 2218 20/2013/DRA (fin whale), 34/2014/DRA (fin whale), 76/2007/DRA (sei whale) (**M.A. Silva**).
- 2219 Short-finned pilot whales were tagged under authorization from NMFS. Bottlenose dolphin
- tagging was conducted under NMFS Scientific Research Permit No. 15543 and approved by
- 2221 Mote Marine Laboratory's IACUC (**R.S.W.**).
- 2222
- 2223 Fishes
- 2224 Tagging procedures were approved by the Committee on Ethics for the Use of Animals of the
- 2225 Universidade Federal Rural de Pernambuco (CEUA #23082.009679/2009 and
- 2226 #23082.025519/2014). Work permits granted by the Instituto Chico Mendes para a Conservação
- 2227 da Biodiversidade (ICMBio #43305–6 and #15083-8) (A.S.A.).
- 2228 Tagging in the Philippines was performed in collaboration with the respective Regional Offices
- 2229 of the Department of Environment and Natural Resources, the Department of Agriculture-Bureau
- 2230 of Fisheries and Aquatic Resources and the Palawan Council for Sustainable Development
- 2231 (Wildlife Gratuitous Permit 2017-13). All research in Tubbataha Reefs Natural Park was done in collaboration with the Tubbataha Management Office (C, A)
- 2232 collaboration with the Tubbataha Management Office (G.A.).
- 2233 Tagging procedures in the Bay of Biscay followed established guidelines that met ethical
- reviews, with scientists limiting handling time and stress as much as possible during attachment(I.A.).
- 2236 Tagging procedures were approved and conducted under Australian Fisheries Management
- Authority Scientific Permit #901193 and Great Barrier Reef Marine Park Authority G11/33231.1
- 2238 (A. Barnett).

- 2239 All procedures for whale shark tagging in the Red Sea were approved by the Institutional
- 2240 Biosafety and Bioethics Committee (IBEC) of the King Abdullah University of Science and
- 2241 Technology. KAUST IBEC serves as the registered (HAP-02-J-042) local committee for all
- 2242 National Committee of Bioethics (NCBE)-regulated activities including animal-related research.
- 2243 (M.L. Berumen and J.E.M.C.).
- 2244 Tagging was conducted with the permission of Chico Mendes Institute for Biodiversity
- 2245 Conservation (number 50119-1), of the Brazilian Ministry of the Environment. Shark capture
- and tagging methods were approved by the Commission of Ethics on the Usage of Animals of
- Federal Rural University of Pernambuco (licence number 054/2013, protocol number
 23082.022567/2012) (N.P.A.B.).
- 2249 Tagging procedures were approved by Stanford University IACUC, the National Oceanic and
- Atmospheric Administration (NOAA), and the California Department of Fish and Wildlife (**B.A.B.**).
- 2252 Tagging of blue, porbeagle and shortfin make sharks in the northwest Atlantic was conducted in
- accordance with the animal care guidelines of DFO and the Canadian Council on Animal Care (S.E.C.).
- 2255 Tagging was conducted with approval by the Province Sud of New Caledonia under permit
- 2256 6024-4916/DENV/SMer and authorization issued by Affaires Maritimes for Chesterfield field 2257 trips (C110-3510-263/MM) (**E.E.G.C.**).
- Tagging was conducted with approval by South African Institute for Aquatic Biodiversity
 Animal Ethics (Ref#25/4/1/7/5_2019-04) (**R.D.**).
- Whale sharks in Madagascar were tagged by Centre National de Recherches Océano- graphiques
 (CNRO) in July 2016 under permit number No 16-12-CNRO-N (S. Diamant).
- Tagging was conducted under permit from the St Helena Government (SHG 20-SRE-01)(A.D.M.D.).
- 2264 Blue sharks (tagged in Irish waters) were tagged under license AE191130/I007 AE19130/P002
- and issued by the Irish Health Products Regulatory Authority (HPRA) and complied with the EU
 Directive 2010/63/EU for scientific research on animals (T.K.D.).
- 2267 Manta ray tagging procedures were approved by the Raja Ampat Marine Protected Area
- 2268 Management Authority and were in accordance with the protocols established by Conservation
- 2269 International Indonesia's and University of Auckland's Animal Ethics Committees (University of
- 2270 Auckland AEC approval #002228). Whale shark tagging was conducted under permits issued by
- the Cendrawasih Bay National Park Authority (SIMAKSI SI.18/BBTNTC-2/TEK/2015,
- 2272 SIMAKSI SI.46/BBTNTC-2/TEK/2015, and SIMAKSI SI.05/BBTNTC-2/TEK/2016). Tagging
- 2273 procedures were approved by the Cenderawasih Bay National Park Authority and are in
- accordance with the protocols established by Conservation International Indonesia's animal
- 2275 ethics review committee (A. Sianipar, E.S. and M.V.E.).
- 2276 Great white sharks were tagged in New Zealand waters according to the protocols specified in
- 2277 Department of Conservation Animal Ethics Committee approvals AEC278, AEC216 and
- AEC260. Mako and porbeagle sharks were tagged according to the code of practice for ethical
- 2279 conduct of tagging carried out by the National Institute of Water and Atmospheric Research
- 2280 (NIWA Animal Ethics Committee 2009) (M.P.F., B.F. and C.A.D.).

- 2281 Tagging was approved by Griffith University ethics (ENV/16/08/AEC) and Ocean and Coast
- 2282 Research animal ethics approval (CA 2010/11/482), with fieldwork conducted under permits
- 2283 6024-4916/DENV/SMer (New Caledonia), G10 33187.2 (Great Barrier Reef Marine Park
- Authority), 143005 (Queensland Fisheries), QS2010 GS065 (Great Sandy Marine Park) and
- 2285 LHIMP/R/2012/009 (Lord Howe Island) (J.G. and J.M.W.).
- 2286Tagging was conducted under permit MAF/LIA/22 to conduct scientific marine animal research2287supplied by the Department of Marine Resources, Bahamas to Bimini Biological Field Station
- 2288 Foundation (**T.L.G.**).
- 2289 Tagging was conducted under permits from the NMFS Highly Migratory Species Division and
- under the University of Miami IACUC. Additionally, blacktip shark tagging was conducted
 under permits from Florida Fish and Wildlife, Everglades National Parks; bull shark tagging was
- 2291 conducted under permits from the Florida Keys National Marine Sanctuary, Florida Fish and
- 2293 Wildlife, and the Biscayne and Everglades National Parks; and great hammerhead shark and
- tiger shark tagging was conducted under permits from the Florida Keys National Marine
- 2295 Sanctuary, Florida Fish and Wildlife, Bahamas Department of Marine Resources, and the
- 2296 Biscayne and Everglades National Parks (N.H.).
- Tagging procedures were conducted under Galapagos National Park Permits PC-13-01, PC-3711. PC-01-14, PC-51-15, PC-69-16, PC-34-17, and MAE-PNG/CDS-2012-0020. Field methods
 were also approved under University of California, Davis IACUC #16022 (A.R.H.).
- Tagging procedures were approved by the University of Windsor Animal Care Committee with a
 permit through Coastal Oceans Research and Development Indian Ocean (CORDIO) (N.E.
 Hussey).
- 2303 Tagging was conducted under Flinders University Animal Welfare Ethics Permits E349 and
- E360, and was authorised by the Victorian Department of Primary Industries under General
- 2305 Research Permit RP1048 and PIRSA Ministerial Exemptions Section 115: 9902064 and 9902094
- 2306 (C.H.).
- 2307 Tagging procedures for scalloped hammerhead and Galapagos sharks were approved by the
- 2308 Zoological Society of London's ethics committee under the project code BPE/0708. Research
- 2309 tagging activities around Mikomoto Island, Japan, were communicated to and approved by
- fisheries officers within the Japanese government (a formal research permit was not required)(D.M.P.J.).
- 2312 For South African white sharks, all research methods were approved and conducted under the
- 2313 South African Department of Environmental Affairs: Oceans and Coasts permitting authority
- 2314 (Permit #RES2012/OCEARCH/umbrella-project) (A.A.K.).
- Tagging was conducted with the full approval of the Instituto Chico Mendes de Conservação da
 Biodiversidade of the Brazilian Ministry of the Environment (permit no. 14124) (B.C.L.M.).
- 2317 Tagging procedures were reviewed and approved by the Seychelles Bureau of Standards, the
- 2318 Seychelles Ministry of Environment, Energy and Climate Change, and The University of
- 2319 Western Australia (RA/3/100/1480) (L.R.P.).
- 2320 Tagging was conducted under Direcção-Geral de Alimentação e Veterinária ethics approvals
- from Decreto-lei N° 129/92 (6 de julho); Portaria N° 1005/92 (23 de outubro) (**N.Q.**).

- 2322 Tagging was carried out under the general auspices of Consejo Nacional de Ciencia y Tecnología
- 2323 (CONACYT), Dirección General de Vida Silvestre (DGVS), Secretaría del Medio Ambiente y
- 2324 Recursos Naturales (SEMARNAT), and Comisión Natural de Áreas Naturales Protegidas
- 2325 (CONANP). These are the relevant Mexican authorities governing all research actions on
- 2326 wildlife and protected animals and areas in Mexico. CONACYT registration: RENIECYT No.
- 2327 030 (currently 1602199) and 13920. DGVS authorization numbers are: SGPA/DGVS/02677/08,
- 2328 SGPA/DGVS/02888/09, SGPA/DGVS/03848/10, SGPA/DGVS/03155/11,
- 2329 SGPA/DGVS/03362/12, SGPA/DGVS/05555/16 and SGPA/DGVS/05970/17 (**D.R.**).
- 2330 All tagging was conducted under animal ethics approvals from Murdoch University's Animal
- Ethics Committee (permit numbers: W2058/7; W2402/11; R2926/17) and an animal ethics
- 2332 permit from The University of Queensland: SBS/085/18/WA/INTERNATIONAL. Permits to
- 2333 conduct research on wildlife in Western Australia were issued by the Western Australian
- 2334 Department of Environment and Conservation (DEC) (permit numbers: SF007471; SF007949;
- 2335 SF008572) and Department of Parks and Wildlife (DPaW) (permit numbers: SF009184;
- 2336 SF009897; SF010414; SF010781; 08-000533-2; 08-002082-2) (**S.D.R.**).
- 2337 Tagging was conducted with permission by the Qatar Ministry of Environment (**D.P.R.**).
- 2338 Tagging in Mozambique was compliant with ethics guidelines from the University of
- 2339 Queensland's Animal Ethics Committee and was conducted under their approval certificate
- 2340 GPEM/186/10/MMF/WCS/SF. Madagascan fieldwork was conducted with the approval of and
- 2341 in partnership with the CNRO in Madagascar. Filipino fieldwork was performed in collaboration
- with the respective Regional Offices of the Department of Environment and Natural Resources,
- 2343 the Department of Agriculture-Bureau of Fisheries and Aquatic Resources and the Palawan
- 2344 Council for Sustainable Development (Wildlife Gratuitous Permit 2017-13) (C.A.R.).
- Tagging methods for broadnose sevengill sharks (*Notorynchus cepedianus*) were approved by the University of Tasmania Animal Ethics Committee (Approval No A0011590) (**J.M.S.**).
- 2347 Tagging procedures were approved by the Marine Biological Association of the UK (MBA)
- 2348 Animal Welfare Ethical Review Body (AWERB) and licensed by the UK Home Office through
- 2349 Personal and Project Licences under the Animals (Scientific Procedures) Act 1986 (D.W.S.).
- 2350 Smooth hammerhead shark (Sphyrna zygaena) tagging was approved by the Massachusetts
- 2351 Division of Marine Fisheries. Porbeagle shark tagging was approved by the University of
- 2352 Massachusetts, Dartmouth IACUC (Protocol #05-07). White shark tagging was conducted under
- 2353 Exempted Fishing Permits (SHK-EFP-11-04, SHK-EFP-12-08, SHK-EFP-13-01, SHK-EFP-14-
- 2354 03) issued to the Massachusetts Division of Marine Fisheries by the NMFS Highly Migratory
- 2355 Species Management Division (G. Skomal).
- Tagging procedures were approved by the University of California, San Diego IACUC (protocol
 S12116) (J.D. Stewart).
- 2358 Whale shark tagging procedures were approved by the University of Western Australia
- (RA/3/100/1110; RA/3/100/1437), University of Adelaide (S-2009-109), or Charles Darwin
 University Animal Ethics Committees (M.T. and M.G.M.).
- 2361 Tagging data according to protocols approved by the South African Department of
- 2362 Environmental Affairs: Oceans and Coasts (now the Department of Forestry, Fisheries and the
- 2363 Environment) and adhered to the legal requirements of South Africa. All research methods were

- approved and conducted under the South African Department of Environmental Affairs: Oceans
- and Coasts permitting authority (Permit #RES2012/OCEARCH/KOCK) (A. Towner).
- Tagging procedures were approved by the Nova Southeastern University IACUC (#064-398-150203) (B.M.W.).
- 2368 Tagging was conducted under permits given by the Subsecreataría de Pesca y Acuicultura de
- 2369 Chile. Resolución exenta (Undersecretary of Fishing and Aquaculture) (P.M.Z.).
- 2370
- 2371 Penguins
- 2372

2373 Tagging procedures for little penguins from Montague Island were approved by the Macquarie

- 2374 University Animal Ethics Committee (Animal Research Authority2014/057), and work was
- conducted under Office of Environment and Heritage NSW Scientific Licence SL100746 (G.C.and R.H.).
- 2377 Tagging procedures were conducted under approval from Monash University Animal Ethics
- 2378 Committee (approval numbers BSCI/2006/12, BSCI/2010/22, BSCI/2011/33), Phillip Island
- Animal Experimentation Ethics Committee (approval numbers 3.2007, 2.2010, 3.2011, 2.2014,
- 2380 7.2017), and research permit issued by the Department of Sustainability and Environment of
- Victoria, Australia (permit numbers 10003848, 10004360, 10005601, 10005605, 10006148, 10007320, 10008506) (A. Chiaradia).
- 2383 Tagging procedures on little penguins off South Australia, were conducted under approval by the
- 2384 South Australian Department of Primary Industries and Regions (PIRSA) Animal Ethics
- 2385 Committee (32-12), and Department for Environment and Water (DEW) (Scientific Permit
- 2386 A24684) (**S.D.G.**).
- Tagging procedures were approved by the Australian Animal Ethics Committee (Department for the Environment and Heritage) and the University of Tasmania Animal Ethics Committee Work was carried out under Macquarie Island special permits M1/3/95 and MI/13/96 (**M.A.H.**).
- Tagging procedures were permitted under US Antarctic Conservation Act Permits (Permit
 #2017-012). Field protocols were approved by the University of California San Diego IACUC
- 2392 (S05480) (data used courtesy of Jefferson T. Hinke).
- 2393 Adelie penguin (*Pygoscelis adeliae*) tagging procedures were approved by the TAAF ethic
- committee and the French regional ethic committee. King penguin (Aptenodytes patagonicus)
- handling procedures were approved by the Ethical Committee of the French Polar Institute
- 2396 (Institut Polaire Paul-Emile Victor). Authorizations to enter the king penguin breeding site
- 2397 (permits nos. 2005–191, 2006–67) and handle birds (permits nos. 99/346/AUT, 00/240/AUT,
- 2398 01/315/AUT, 01/322/AUT, 2003–113, 2003–114, 2004–182, 2004–183, 2005–203 and 2006–73)
- were delivered by the French Ministère de l'Aménagement du Territoire et de l'Environnement
 (MATE) and TAAF (Y.R.).
- Animal handling procedures were approved by the joint University of Cambridge / British
- 2402 Antarctic Survey Animal Ethics Committee (**P.N.T.**).
- 2403 Tagging procedures were approved by the Animal Ethics Committee of the Australian Antarctic
- 2404 Division (ATEP-12-13-4086-4088-SUMMER) (**B.W.**).
- 2405

- 2406 Polar bears
- 2407 Tagging procedures were conducted under USFWS research permit MA 690038 Animal Care
- and Use Committees of the US Geological Survey (assurance no. 2010–3) (A.M.P.). 2409
- 2410 Seals
- 2411 Tagging was permitted by the Russian Federal Veterinary and Agricultural Control Service
- 2412 (Rosselkhoznadzor, Kamchatka and Koryakia regions, Permit No. 1194) and was approved by
- 2413 the Alaska Sea Life Center IACUC (**R.D.A.**).
- Tagging procedures were approved by the Adelaide University Animal Ethics (permit S80-2004) and South Australia Department for Environment and Heritage (permit A24684-3) (**A.M.M.B.**).
- 2416 Weddell seals (Leptonychotes weddellii) tagged in Dumont d'Urville, Adélie Land by LOCEAN
- 2417 laboratory were treated in accordance with the Institut Paul-Emile Victor (IPEV) ethical and
- 2418 Polar Environment Committees guidelines (J. Charrassin).
- 2419 Animal use protocols for northern elephant seal tagging was reviewed and approved by the
- 2420 University of California at Santa Cruz IACUC and followed the guidelines established by the
- 2421 ethics committee of the Society of Marine Mammalogy. Research was carried out under NMFS
- 2422 permits: #786-1463 and #87-143. Southern elephant seal (*Mirounga leonina*) captures were
- conducted under NMFS permit No. 87-1851-00. All animal procedures were approved by the
- 2424 IACUC at University of California Santa Cruz. Weddell seal handling protocols were approved
- 2425 by the University of Alaska Anchorage and University of California Santa Cruz's IACUCs.
- 2426 Research and sample import to the United States were authorized under the Marine Mammal
- permit No. 87-1851-04 issued by the Office of Protected Resources, NMFS. Research activities
- on southern elephant seals and Weddell seals were also approved through Antarctic Conservation Act permits while at McMurdo Station ($\mathbf{D} \mathbf{P} \mathbf{C}$ and $\mathbf{P} \mathbf{W} \mathbf{P}$)
- 2429 Act permits while at McMurdo Station (**D.P.C.** and **P.W.R.**).
- 2430 Ringed seal (Pusa hispida) handling and tagging was approved by the University of Windsor
- Animal Care Committee (AUPP #12-12,13-10) and a DFO License to Fish for Scientific
- 2432 Purposes (S-12/13-1019-NU) (S.H.F. and D.J.Y.).
- 2433 Australian sea lion (Neophoca cinerea) tagging procedures were approved by the PIRSA Animal
- 2434 Ethics Committee (32-12), South Australian DEW (Scientific Permit A24684), and Western
- 2435 Australian Department of Environment and Conservation (Licence to Take Fauna for Scientific
- 2436 Purposes SF009529). Long-nosed fur seal tagging procedures were approved by the PIRSA
- 2437 Animal Ethics Committee (32-12) South Australian DEW, Scientific (Permit A24684) (S.D.G.).
- 2438 Tagging procedures for Australian fur seal (Arctocephalus pusillus doriferus) and New Zealand
- fur seal (*Arctocephalus forsteri*) were approved by the Macquarie University Animal Ethics
- 2440 Committee (Animal Research Authority2014/057), and work conducted under Office of
- 2441 Environment and Heritage NSW Scientific Licence SL100746. Weddell seal tagging procedures
- were approved by Macquarie University (#3223) ARA 2014_057 (**R.H.**) or approved by the
- 2443 New Zealand Department of Conservation, Ministry of Foreign Affairs and Trade, and NIWA
- 2444 Animal Ethics Panel (DOC-69331-MAR) (M.P.).
- 2445 Animal Ethics were obtained from NIWA to manipulate New Zealand sea lions (Phocarctos
- 2446 *hookeri*) at Campbell Island, with the proviso that all work was undertaken with approval from
- the Department of Conservation and the NZ Department of Conservation permit issued under the

- 2448 Marine Mammal Protection Act (1978). Southern elephant seal tagging procedures were 2449 approved by University of Tasmania Animal Ethics (permit A0014523) (**M.A.H.**).
- All animal captures and procedures were authorised under NMFS permits (numbers 87-1593 and
- 2451 87-1851-00) and approved by the University of California, Santa Cruz IACUC. Fieldwork in
- 2452 Antarctica was approved by the Antarctic Conservation Act (L.A. Huckstadt).
- Tagging procedures were approved by the UCC Animal Ethics Committee, Irish National Parks
 & Wildlife Service, and HPRA (M.J.).
- 2455 Southern elephant seals, Antarctic fur seals (*Arctocephalus gazella*), Weddell seals, crabeater
- 2456 seals (Lobodon carcinophaga) and leopard seals (Hydrurga leptonyx) were tagged under ethics
- and permits provided by the Brazilian Antarctic Programme "in lieu" of SCAR as their local
- 2458 representatives for all the field work conducted on pinnipeds at Elephant Island, South Shetlands
- 2459 (**M.M.C.M.**).
- 2460 Tagging procedures were conducted under the permit #572/208 approved by the National
- Administration of Aquatic Resources, Ministry of Livestock, Agriculture and Fisheries (DINARA), Uruguay (F C R)
- 2462 (DINARA), Uruguay (**F.G.R.**).
- 2463 Tagging procedures were approved by the Dirección Nacional del Antártico, Buenos Aires,
- 2464 Argentina, and were carried out according to the Scientific Committee on Antarctic Research
- 2465 Code of Conduct for Animal Experiments under University of New South Wales Animal Care
- and Ethics Committee (Protocols 08/103B and 11/112A), and the Animal Care and Ethics
- 2467 Committee of the Antarctic Science Advisory Committee (permit number 1144) (**T.L.R.**).
- 2468 Northern fur seal (*Callorhinus ursinus*) tagging in the Pacific North East and Pacific East Central
- 2469 was conducted in accordance with and under the authority of the United States Marine Mammal
- 2470 Protection Act (NMFS Permits 782–1455 and 782–1708). At the time this work was conducted
- 2471 there was no additional requirement for review of these procedures by an institutional review
- board or ethics committee. In 2010, a NMFS IACUC was established for the Alaska Fisheries
- and Northwest Fisheries Science Centers and the capture and handling protocols were reviewed
- and approved by this committee (J.T.S. and R.R.).
- 2475 Harbor seal (*Phoca vitulina*) studies in Scotland were carried out under UK Home Office licence
- under the Animal (Scientific Procedures) Act 1986 (PIL nos. 60/3303, 60/4009 and 70/7806),
- following approval by the University of St Andrews animal welfare and ethics committee.
- 2478 Licences to capture and release animals in the wild for research were also granted by Marine
- 2479 Scotland Licensing (**P.M.T.**).
- 2480 Animal handling and instrumentation complied with animal care regulations and applicable
- 2481 national laws of Ecuador. This research was approved by the Chancellor's Animal Research
- 2482 Committee at University of California, Santa Cruz. The appropriate animal use and care
- 2483 committee of Ecuador (Parque Nacional Galapagos) approved all research protocols. This work
- was performed under the permit No PC-11-08 and PC-043-09 and authorization No. 084/06
- 2485 PNG of the National Park service, Galapagos (S.V.).
- 2486 Grey (Halichoerus grypus) and harbor seals were caught under licenses Number 05/475/AUT,
- 2487 05/485/AUT, 06/82/AUT, 07/481/AUT, 08/346/DEROG, 08/347/DEROG, 10/102/DEROG,
- 2488 11/873/DEROG, 11/874/DEROG, and 13/422/DEROG delivered by the French ministry of the 2489 environment (C V)
- 2489 environment (C.V.).

- 2490 California sea lion (Zalophus californianus) capture and procedures were approved by the
- 2491 University of California Santa Cruz Chancellor's Animal Research Committee (CARC) protocol
- 2492 (COST 01.10) and authorized under National Marine Fisheries Service permit number 87-1593-
- 2493 05 (**M.J. Weise**)
- 2494
- 2495 Sirenians
- 2496 Dugong tagging was conducted under the conditions of ethics permit DEC AEC 2009/112497 (R.A.C.).
- Permits required to capture and satellite track dugongs were obtained from the James Cook
 University Animal Ethics Committee (Permits A1735 and A1936) and the North (609121552013/JJC) and South (3157- 2012/ARR/DENV) Provinces of New Caledonia (C.C.).
- 2501 Tagging procedures were approved by the Charles Darwin University Animal Ethics Committee
- and wildlife research permits were obtained from the Parks and Wildlife Commission of the
- 2503 Northern Territory (S.D.W.).
- 2504 Manatee tagging procedures were carried out in accordance with the USFWS Permits
- 2505 MA107933-1 and MA37808A-0, Alabama Department of Conservation and Natural Resources,
- and Alabama Division of Wildlife and Freshwater Fisheries annual permits. Approvals obtained
- by the University of South Alabama IACUC for protocols 581568 and 1038636 (**R.H.C.**).
- 2508
- 2509 Turtles
- 2510 Tagging was conducted under permits from Dirección General del Medi Natural de la
- 2511 Generalitat Valenciana, Generalitat de Catalunya, Consejeria de Medio Ambiente y Ordenación
- del Territorio de la Junta de Andalucia, and Región de Murcia. A general permit for tagging
- 2513 adult females was obtained from Ministerio para la Transición Ecológica y el Reto Demográfico
- 2514 (GPM/BDM/AUTSPP/23/2020) (S.A. and E.J. Belda).
- 2515 The Indonesian Institute of Sciences provided research permits for telemetry deployments at the
- 2516 nesting beaches. Telemetry deployments at California foraging grounds were conducted under
- 2517 Endangered Species Act permit nos. 1159, 1227, and 1596 (S.R.B.).
- Queensland Scientific purposes permit and a University of Queensland Animal Ethics permit(H.A.C.).
- 2520 Tagging procedures were conducted under permits granted by the Commonwealth of Dominica
- 2521 Ministry of Agriculture and Forestry to Domenicia's Sea Turtle Conservation Organisation Inc2522 (R.W.C.).
- 2523 Green turtle (Chelonia mydas) tagging procedures were carried out in compliance with Mexican
- regulations (permit SGPADGVS/SEMARNAT, Mexico, No.09583/15). Hawksbill turtle
- 2525 (Eretmochelys imbricata) tagging was carried out in compliance with Mexican regulations
- 2526 (permit SGPADGVS/SEMARNAT Mexico, No.09583/15) (E. Cuevas-Flores).
- 2527 Loggerhead turtles (*Caretta caretta*) were handled under license "N° 04/IFCN/2018- FAU
- MAO" and previous licenses issued by the Government of the Autonomous Region of Madeira (T.D.).
- 2530 Leatherback turtle tagging procedures were conducted under NMFS Endangered Species Act
- 2531 Section 10 Permits #1557 and #15672, University of New Hampshire IACUC #060501 and

- 2532 #090402, and University of Massachusetts IACUC #2010-0019. Turtle disentanglement was 2533 conducted under the authority of NOAA 50 CFR Part 222.310 (K.L.D.).
- 2534 All sea turtle research was conducted under NMFS Permit 1260 and 16733 to take protected
- 2535 species for scientific purposes and USFWS permits TE-676379-4 and TE676379-5 issued to the
- 2536 NMFS Southeast Fisheries Science Centre (SEFSC) and according to IACUC-reviewed
- 2537 procedures outlined in the NMFS SEFSC Sea Turtle Research Techniques Manual (L.L.D.).
- 2538 Loggerhead and green turtle tagging procedures were conducted under permit issued by the
- 2539 wildlife agencies of Buenos Aires and Río Negro provinces and the National Wildlife Agency of
- 2540 Argentina (V.G.C.).
- Green turtle tagging procedures were conducted within the Statia National Marine Park 2541
- 2542 programme and complied with all relevant national legislation. Hawksbill turtle tagging was
- 2543 conducted within the Statia National Marine Park and St Maarten Marine Park programmes and 2544 complied with all relevant national legislation (N.E.).
- 2545 Leatherback turtle tagging procedures were reviewed by the University of New Hampshire 2546 IACUC (060501) (B.J.G.).
- 2547 Leatherback turtle were tagged under permit number SGPA/DGVS/08562/17. Green and
- 2548 hawksbill turtle tagging procedures were authorized by the SEMARNAT (permit numbers
- 150496-213-03, 280597-213-03, 190698-213-03, 280499-213-03, SGPA/DGVS/002m 2549 2550 SGPA/DGVS/05137/12, SGPA/GDVS/02259/14, and SGPA/DGVS/04478/15) (C.E.H.).
- 2551 Tagging procedures were approved by Swansea University and Deakin University Ethics
- 2552 Committees and the British Indian Ocean Territory (BIOT) Administration of the UK Foreign
- 2553 and Commonwealth Office. Research was endorsed through research permits (0002SE12,
- 2554 0007SE15, 0002SE17, 0006SE18) from the Commissioner for BIOT and research complied with
- 2555 all relevant local and national legislation (G.C.H.).
- 2556 Sea turtle tagging procedures and fieldwork were directly approved by the Centro
- TAMAR/IBAMA/ICMBio. Fundação ProjetoTAMAR has MMA/IICMBio/SISBIO Nº 42760 2557 2558
- permit (P.H.L. and E.A.P.S.).
- 2559 Tagging procedures for rehabilitated loggerhead sea turtles were conducted under the
- 2560 authorization of blanket permit from USFWS to NOAA NMFS. Loggerhead sea turtles acquired 2561 via capture or incidental capture were taken under the authority of NMFS Research permit 16134 2562 (G.L.).
- 2563 Green turtle tagging procedures were conducted with permission from the Administrator of
- 2564 Ascension Island. Leatherback turtle tagging was conducted under permits from Ezemvelo
- 2565 KwaZulu Natal Wildlife. Loggerhead turtle tagging was conducted with approval from the
- 2566 ethical committee of the University of Pisa (P.L.).
- 2567 Tagging procedures were authorized under the Peru Instituto Nacional de Recursos Naturales
- 2568 (INRENA) permits 015-2002-INRENA-J-DGFFS-DCB, 070-2003-INRENA-IFFS-DCB, 068-
- 2569 2004-INRENA-IFFS-DCB, 025-2005-INRENA-IFFS-DCB and 002-2006-INRENA-IFFS-DCB
- 2570 (**J.C.M.**).
- 2571 Tagging operations were authorized by the Dirección General de Sostenibilidad de la Costa y del
- Mar (Ref DIV/BDM/AUTSSP/58/2015, Spanish Government) (D.M.). 2572

- 2573 Green turtle tagging was conducted under permits of NOAA, Federated States of Micronesia,
- and Republic of Marshall Islands. Hawksbill turtle tagging was conducted under permits from
- 2575 NOAA, the USFWS, the Hawai'i Division of State Parks, and the Mexico Comisión Nacional de
- 2576 Áreas Naturales Protegidas (**D.M. Parker**).
- 2577 Loggerhead turtles tagging procedures off of the Baja California Peninsula, Mexico, were
- 2578 conducted in full compliance with CARC/IACUC protocol at UC Santa Cruz and research was
- authorized by the Mexican government through SEMARNAP and SEMARNAT permits
- 2580 150496-213-03, 280597-213-03, 190698-213-03, 280499-213-03, 280700-213-03,
- 2581 SGPA/DGVS/002 4661, SGPA/DGVS/10358, and SGPA/DGVS/03501/06 (H.P.).
- 2582 Tagging procedures were authorised by the Environment Agency Abu Dhabi, the Environment
- 2583 & Protected Areas Authority, Sharjah, the Environment Studies Center at Qatar University, the
- 2584 Qatar Ministry of Environment, the Oman Ministry of Environment and Climate Affairs, and the 2585 Department of Environment, Iran (**N.J.P.**).
- 2586 Leatherback turtles tagging procedures were conducted under licence (# 27/01 and 73/08) from
- 2587 the Fauna Department-Ministry of Cattle, Agriculture and Fishing of Uruguay (L.P. and M.

2588 Lopez Mendilaharsu).

- Tagging permissions were given by Oman's Ministry for Regional Municipalities, Environment
 and Water Resources (A.F.R.).
- Tagging was performed with the permit of the Environmental Ministry of the DominicanRepublic Government (J. Tomás).
- 2593 Permissions for sea turtle rehabilitation work were given by the Dubai Wildlife Protection Office2594 (D.P.R.).
- 2595 Tagging procedures were conducted under approval from the National Marine Park of Zakynthos
- 2596 (permits from 2000–2012), the Animal Ethics Committee of Deakin University (B0X2015-17),
- and the Greek Ministry of Environment (Permit: 151503/162) (G. Schofield).
- 2598 Tagging procedures were conducted under approval from the Dakshin Foundation Animal
- 2599 Research Ethics Review Committee. In the Andaman and Nicobar Islands, permits were issued
- 2600 to tag ten leatherback sea turtles with satellite transmitters from the Ministry of Environment and
- 2601 Forests (Wildlife Division), Government of India, on 16th December 2008 (F.No.1-4/2007 WL-I
- 2602 (pt-1)). Research permits from the Forest Department, Andaman and Nicobar Islands
- 2603 (CWLW/WL/47/393) and other relevant permits from the Andaman and Nicobar Administration
- were also obtained to carry out the field work in Little Andaman Island (K.S.).
- 2605 Tagging procedures were performed in accordance with the Stanford University Protocol for the
- 2606 Care and Use of Laboratory Animals (APLAC no. 13848). The Costa Rican Ministry of Natural
- 2607 Resources and the Environment provided research permits (G.L.S.).
- 2608 Green turtle tagging was conducted under permit approved by the Western Australian
- 2609 Department of Biodiversity, Conservation and Attractions. Tags were deployed by RPS Group -
- 2610 Perth WA (lead by former employee **D.W.**) on behalf of Woodside Energy Group Ltd.
- 2611 Tagging procedures were conducted under permission obtained from the Viceconsejería de
- 2612 Medio Ambiente of the Gobierno de Canarias. Cape Verde did not require permission from the
- 2613 government at that time (N.V.).

- 2614 Hawksbill turtle and olive Ridley turtle (*Lepidochelys olivacea*) tagging procedures were
- 2615 conducted under approval from Charles Darwin University Animal Ethics Committee and
- 2616 wildlife research permits (A4005) from Parks and Wildlife Commission of the Northern
- 2617 Territory (**S.D.W.**).
- 2618 Research protocols for capturing and deploying satellite transmitters on flatback turtles were
- approved by an authorised ethics committee (SA 2015/11/531) and authority under the Nature
- 2620 Conservation Act 1994 (I.B., C.A.M.H., A. Barnett, N.E.W.).

2621 Tracking data collection and processing

2622 Tagging devices were deployed across more than three decades from 1985 to 2018 around the 2623 global ocean, resulting in a total of almost 11 million positions (after data curation: 6,854,440 2624 positions) collected with different sensor systems and technologies for transmitting data. These 2625 included tagging devices using the Argos doppler-shift localization system (argos-system.org), 2626 GPS (global positioning system) and Fastloc GPS, as well as light-level geolocation tags (also 2627 termed global location sensor; GLS). Animals within taxa were captured (or tagged remotely) by different teams using a range of methods after the responsible team leader obtained all licenses 2628 and ethical permissions (see "Fieldwork and Data Collection"). All birds including penguins 2629 2630 were mostly caught at nest sites using poles, traps, or nets, and tags generally attached dorsally or to a leg. Most cetaceans were tagged from the research vessel using crossbows, air-powered 2631 2632 systems or poles to get tags attached to the dorsal fin or its vicinity. Fishes were mostly captured 2633 with baited hooks or purse-seine nets and tags typically attached to the first dorsal fin using a 2634 tether affixed to a dart or by fixing it with stainless steel bolts. Satellite collars were used for 2635 polar bears after immobilisation using rapid-injection darts. Seals were mostly captured with nets 2636 and sedated before tag deployment on the head or along the dorsal midline. All sirenians were tagged using a peduncle belt linked to the tag by a tether. Most turtles were captured at nesting 2637 2638 beaches or at sea using nets or the by-hand 'rodeo' technique and tags glued to their carapace, 2639 except for leatherback turtles (Dermochelys coriacea) for which a harness ("backpack"), towed-2640 tag or surgical techniques were used for attachment. All animal handling and tagging procedures were completed by trained personnel under permissions granted by ethical review bodies and in 2641 2642 accordance with all relevant ethical regulations in the jurisdictions in which they were performed 2643 with specific approvals obtained by each data owner who was individually responsible for 2644 adhering to regulations and supervision of all procedures (details provided in "Animal Ethics 2645 Information").

2646 Tracking datasets were collated after a lead author (representing each tagging research team) 2647 provided three csv files, each including species metadata, tracking data, and the team description. 2648 All datasets were requested with the least amount of processing possible, with all Argos, GPS 2649 and fastloc GPS data (~90% of the tracking data) provided as 'raw' position estimates. GLS 2650 positional data (for some birds and fishes only) were provided after estimation of longitude and 2651 latitude from the ambient variables recorded in the device (i.e., light intensity and elapsed time, 2652 but also depth and temperature for fishes). For birds, GLS position estimates provided were 2653 obtained in two ways: (i) through the Geolight package(112) in R(113) after carrying out a pre-2654 and a post-calibration (seven days) to estimate an average value for the sun elevation parameter 2655 needed for calculations, or (*ii*) through the BASTrack software suite (British Antarctic Survey) 2656 after identifying sunrise and sunset times based on light curve thresholds and with longitude and 2657 latitude calculated from the time of local midday and day length, respectively. The exception was 2658 for the dataset for the hybrid complex of three *Pterodroma* species(114), referred here to as 2659 Trindade Petrel (Pterodroma arminjoniana), for which the GLS positions were processed with 2660 the R package *TripEstimation(115)*. For fishes, GLS tracks were obtained using pop-up satellite 2661 archival transmitters (PSAT) through satellite-relayed data or archived data from tags physically 2662 recovered. Positions were obtained after data decoding using software provided by the 2663 manufacturers (e.g., Wildlife Computers), where, similarly to bird data, longitude and latitude 2664 are calculated from estimated local time of midnight or midday and day-length, respectively. 2665 These PSAT GLS tracks were further processed with a continuous-time correlated random walk 2666 (CTCRW) Kalman filter using the crawl package(116) in R to produce daily positions, after

2667 filtering with the unscented Kalman filter using sea surface temperature through the UKFSST

2668 package in R and applying a bathymetric correction using the analyzepsat R add-on. PSAT data

- also included shark tracking data from the Tagging of Pacific Predators (TOPP) program which
- 2670 were downloaded from the Animal Tracking Network (ATN) hosted by the Integrated Ocean
- 2671 Observing System (ioos.noaa.gov/project/atn/, downloaded September 2017) for integration in 2672 the Global Shark Movement Project (GSMP; globalsharkmovement.org). These data obtained
- 2673 through GSMP were processed as detailed in (62) to determine daily position data.
- 2674 All data were checked for quality-control and to standardise formats of the multiple and disparate 2675 datasets received. Poor data quality, lack of metadata, misidentified or incomplete tracks, 2676 repeated tracks, or unsolicited processing before data submission (e.g., interpolation of Argos tracked positions) led to the exclusion of 3,051 tracks from 10 species prior to analyses. All 2677 2678 datasets were run through a speed filter using the SDA filter package in R to remove outlier positions. Speeds used ranged for species between $5.4 - 35 \text{ m.s}^{-1} (20 - 126 \text{ km.h}^{-1})$ for birds, 1.6 2679 $-7 \text{ m.s}^{-1} (5.8 - 25 \text{ km.h}^{-1})$ for cetaceans, $0.5 - 11.9 \text{ m.s}^{-1} (1.9 - 42.8 \text{ km.h}^{-1})$ for fishes, 2.1 - 4.22680 $m.s^{-1}(7.5 - 15 \text{ km.h}^{-1})$ for penguins, 0.75 $m.s^{-1}(2.7 \text{ km.h}^{-1})$ for polar bears (but see (117)), 2.0 -2681 10.3 m.s^{-1} (7.2 – 37 km.h⁻¹) for seals, $1.1 - 2.8 \text{ m.s}^{-1}$ (4.1 – 10 km.h⁻¹) for sirenians, and 1.4 - 2.82682 2683 $m.s^{-1}$ (5 – 10 km.h⁻¹) for turtles (refer to table S3 for details, and also for general morphometric 2684 data per species). During this procedure, all Argos data resulting from unsuccessful satellite 2685 uplinks (i.e., with location class Z) were removed from the dataset, keeping only location classes 2686 B, A, 0, 1, 2, and 3, which have increasing accuracy from ~ 160 km to 0.3 km (118). Visual inspection led to further removal of unrealistic GLS locations for some bird species (e.g., 2687 longitude $< 43^{\circ}$ W or $> 98^{\circ}$ W, latitude $< 8^{\circ}$ N or $> 73^{\circ}$ N for Arctic herring gull – *Larus* 2688 2689 smithsonianus). A land mask was applied to all data using the rworldmap package in R and all 2690 locations assigned to land were excluded from analyses. We created 1° grid-cells for all area 2691 included in the world's ocean, and all grid-cells where animal tracking data were not detected 2692 have also been excluded from analyses. Because the area within each grid-cell varies considerably with latitude, all results were calculated based on area following: 2693
- 2694 $A(\theta) = 2\Delta \phi R^2 [\sin(\theta_{max}) \sin(\theta_{min})]$ 2695 where θ is latitude, ϕ is longitude and θ_{max} and θ_{min} are the bounding latitudes of the grid-cell,
- 2696 and R is the average Earth's radius (6,371 km).
- 2697

2698 Addressing tracking data biases

2699 The inherent biases in tracking datasets(63), such as the different data resolution and number of 2700 positions resulting from different devices, higher number of positions commonly obtained 2701 around tagging locations, and different track lengths obtained from devices deployed at the same 2702 time, make analyses challenging. To alleviate some of these potential issues, we gridded data at 2703 1º resolution, keeping only the counts of unique individuals per species. We chose this resolution 2704 because it encompasses most of the known accuracies for most tracking devices, including most 2705 positions obtained by PSAT GLS(62), therefore alleviating most of the effects of position error 2706 estimates on track accuracy. This resolution has also been proposed as the best resolution to use 2707 when performing statistical analyses at large spatial scale(63, 119) or when using 'big data' 2708 approaches(120). To further reduce any potential biases in track accuracy due to the lower 2709 accuracy of GLS data and their limited daily locations (usually only 1 or 2), we repeated all 2710 spatial analyses using only one position per day for each individual, calculated as the centre of

2711 mass of all position estimates obtained per individual in a given day, and also used this dataset 2712 for all time-based calculations. We found these potential methodological biases led to no major 2713 differences in the pattern of results obtained (fig. S5). Furthermore, to avoid overestimating 2714 spatial overlaps due to the 'addition' of locations by interpolation methods – which can lead to 2715 locations being introduced where the animals were likely to have been but which were not 2716 detected by the tracking devices deployed – we considered all the positions that were detected, 2717 rather than interpolating positions for all taxa, except as detailed above (e.g., for the PSAT GLS 2718 daily position data for sharks derived from GSMP). Track interpolations, which are often 2719 calculated between positions up to 20 days apart(61) can result in an additional source of 2720 bias(63) and could inflate our globally important marine megafauna areas. By using only 2721 detected positions (rather than interpolated) and focusing on unique detections for each 2722 individual (instead of number of positions) within each 1º grid-cell, we conservatively estimated 2723 important marine megafauna areas that were also not affected by inflated detections around each 2724 tagging location (i.e., only one position was considered for each individual within 1º resolution 2725 around the tagging location). To further understand the potential effects of the tagging location 2726 bias, we also repeated our spatial analyses after removing all positions around the tagging 2727 location where the probability of finding an individual following a random trajectory from the tagging location was >10%. We did this by estimating the characteristic daily velocity (i.e., the 2728 2729 root mean square displacement, d) for each species, and then using this value to estimate the diffusion constant (D) for a Brownian random walk, as $D = d^2/2T$ (with T = 1 day). We then 2730 compared our curated tracks with those obtained from trajectories generated through a Brownian 2731 2732 random walk with that diffusion constant, when using similar starting locations for each track. 2733 We used these trajectories to estimate the probability of an individual randomly arriving at the 2734 same distance (or further) from each tagging location as that observed in the curated tracks, and 2735 discarded all positions where this probability was >1%. We then used our curated tracks with 2736 new starting positions matching the first location where the probability of randomly being at that 2737 location estimate was <10%, to re-compute our spatial analyses, which resulted in similar 2738 patterns obtained (fig. S6). Finally, to study the effects of spatial resolution on all our results, we 2739 repeated all the analyses at 0.5° and 2° grid-cell resolutions and found similar patterns (see fig. S7, fig. S8). All comparisons were made using the Jaccard similarity coefficient (or Jaccard 2740 index), which is calculated by dividing the size of the intersection of two datasets by the size of 2741 2742 their union, and results in 0 for no intersection between the sets (i.e., complete dissimilarity) and 2743 in 1 for equal sets (i.e., high similarity).

2744

2745 Detection of key movement behaviours

2746 To detect key movement behaviours such as migration (defining migratory corridors) or 2747 residence (potentially indicating feeding, mating or resting areas) throughout the three decades of 2748 tracking data in our multi-taxa global dataset, we used an algorithm based on statistical methods 2749 commonly applied to big data analyses. Our algorithm uses a time series of displacements 2750 calculated as the shortest great-circle distance, i.e., measured along the surface of the sphere, 2751 between two consecutive tracked locations separated by predetermined time-windows (Tw) (as 2752 done in 120 from 1 - 10 days. We then calculated the average displacement per individual and 2753 normalised the displacements by the average displacement per species to account for disparities 2754 in speed across the 111 species considered in our study.

27552756 Detection of migratory corridors

2757 For detecting migratory corridors captured by our tracking dataset, we calculated how coherent the movement direction was within each grid-cell for each species based on the displacements 2758 2759 calculated for Tw = 1 - 10 days. We did this because the results obtained for movement direction 2760 can differ for long- and short-time windows, with the former likely to reflect long term movements in a specific direction (i.e., ignoring potential return trips or other shorter changes in 2761 2762 direction, such as daily trips), and the latter likely to provide displacements that are 2763 unrepresentative of potential migration (i.e., 'noisy' data). We then defined *coherence* (c) per taxon, for each Tw and grid-cell, as the sum of the displacement vectors (w_d) in a particular 2764 2765 direction (i.e., multiplied by the cosine and sine of the angle φ) and then divided by all 2766 displacements in all directions, as:

2767
$$Coherence(T_w, c) = \frac{\sum_{d=1}^{D} w_d \left(\cos \varphi_d, \sin \varphi_d\right)}{\sum_{d=1}^{D} w_d}$$

2768 where D represents all displacements observed in each grid-cell. To scale results across taxa, we

multiplied the average monthly *coherence* by the ratio between the number of grid-cells with observed displacements within each time window (C_{dm}) and the maximum number of grid-cells

2770 observed displacements within each time window (C_{dm}) and the maximized observed over different time windows (max C_{dm}) for each taxon.

2772 The selected taxon-specific displacements calculated for the Tw that resulted in the maximum 2773 number of 1° grid-cells showing coherent movement for that taxon (i.e., 'best Tw'; refer to fig. 2774 S12) were then aggregated at temporal scales of 1, 2, 3, 4, 5, 6 and also 12 months. Considering 2775 multiple temporal scales was necessary due to the differences in movement behaviour across the 2776 many species considered in our study. For example, central place foragers return to start 2777 locations (e.g., colony) in each trip. Using multiple temporal scales is therefore useful to allow detection of movement corridors in both directions avoiding the cancellation of the displacement 2778 2779 vectors occurring in opposite directions (e.g., trip from nest to foraging location cancelled by the 2780 reverse trip). Also, because migratory behaviour is largely unknown or incomplete for many 2781 species (e.g., sharks), we included temporal scales up to 12 months to ensure we captured any previously undetected long-term migration if present. To automate routines, these temporal 2782 2783 scales were programmatically defined considering one month as 365 days/12 (~ 30.4 days). We 2784 then repeated the calculation of *coherence* at each temporal scale for each taxon to find sets of 2785 neighbouring grid-cells where displacements obtained from the tracking dataset indicated 2786 movement in the same direction. We did this by calculating the average direction of all observed 2787 displacements at each grid-cell within each temporal scale (e.g., for a temporal scale of 3 2788 months, we used displacements calculated between 0 and \sim 90 days) and clustering all grid-cells 2789 that resulted in similar average direction (i.e., for which the cosine of the angle between their 2790 directions is > 0.8, i.e., indicating similar direction of movement).

The clustering of grid-cells resulted in a high number of clusters for each taxon and temporal scales. So, we computed the size distribution of clusters of grid-cells with similar average direction for each temporal scale and plotted the cumulative distribution. Then using a Lorenz curve as a parameter-free approach(*121*), we identified the intersection point between the slope of the tangent line at the maximum value (i.e., the slope where the cumulative distribution equals

2796 1) and the *x*-axis in the Lorenz curve plot. This intersection point defines the threshold for
minimum cluster size (i.e., minimum number of 1° grid-cells) defining a movement corridor at
each temporal scale for each taxon (see fig. S13). All clusters with size above the defined
threshold at any temporal scale were considered, and all 1° grid-cells within those clusters were
aggregated and classed as corridors. Because speed is generally expected to be faster while
"migrating", we confirmed speed within resulting corridors was always above average for each
species.

2803

2804 Detection of residence areas

2805 To determine residency-like behaviour indicative of areas where animals might be foraging, 2806 feeding, mating or resting (commonly characterised by slower speeds and greater tortuosity), we 2807 computed the z-scores (dimensionless) of the displacements starting within each grid-cell, 2808 considering the average displacements and respective standard deviations per species for the 2809 'best Tw' identified for each taxon. Each displacement observed in a track belonging to a species 2810 within each grid-cell was assigned a z-score by subtracting the average global displacement of that species from the calculated average displacement and dividing the result by the standard 2811 2812 deviation of the displacements of that species. We then used these values to calculate the average 2813 z-score for each taxon in each grid-cell. If the average z-score calculated within each 1° grid-cell was lower than -1 (i.e., one standard deviation below the average displacement for that taxon), 2814 2815 we considered it as reflecting a residency-like movement behaviour, and the corresponding gridcells were classed as residence area. We used this approach to calculate z-scores across the same 2816 aggregated temporal scales used for detection of migratory corridors (i.e., using sets of 1, 2, 3, 4, 2817 2818 5, 6 and also 12 months) and, for a given taxon, observing residency-like movement behaviour in 2819 any of these scales led to the classification of the grid cell as a residence area for that taxon. To confirm that a random approach to identify areas of residence is not useful, we randomised all 2820 2821 tracks in the dataset by changing the sequence of displacements to break their correlation but 2822 keeping the same start and end point of the trajectories (and therefore the same probability 2823 distribution function) (122). We then repeated the procedure to detect residence areas and see if they would be similar. We then used the Jaccard index (123) to measure the similarity between 2824 each randomised set of residence areas and the original per taxa. Detection of residence areas 2825 2826 was substantially different after track randomisation, confirming space-use by animals was not random (fig. S14). 2827

2828

2829 Statistical Modelling

2830

2831 Input Data

We modelled the probability of finding areas (grid-cells) used as residences or for migration separately for each taxon (except sirenians due to lack of data) using generalised linear models with a binomial error distribution and a logit link function. We develop these models considering

as presences the locations where we have detected the described residence or migratory

2836 behaviours for each taxa and by randomly selecting equal number of locations where tracking

2837 data were available for each taxa but no behaviour was detected (see Table 2).

2839 We then used a total of 13 environmental variables as predictors obtained from various online 2840 datasets (see Supplementary Acknowledgements). The predictors included monthly mean global 2841 sea surface temperatures (sst), ocean surface currents (u and v; respectively, eastward and 2842 northward ocean currents), sea surface height (ssh), salinity (sal), and mixed layer depth (mld) collated from the E.U. Copernicus Marine Service Information (CMEMS) Marine Data Store 2843 (MDS) Global Ocean Physics Reanalysis(124). Dissolved oxygen (O_2) was obtained from the 2844 2845 CMEMS Global Ocean Biology Hindcast replaced in July 2022 by the Global Ocean 2846 Biogeochemistry Hindcast (125). Ocean turbidity (turbidity) and chlorophyll-a concentration 2847 (chla) were obtained from NASA Ocean Biology Processing Group Level-3 SeaWifs (1998-2848 2003) (126) and Modis-Aqua (2003-2018) (127) Ocean Color Data. Atmospheric temperature at 2849 2 m height (*temp2m*) and wind velocity at 10 m height (u10 and v10, respectively representing 2850 eastward and northward direction) were obtained from the European Centre for Medium-Range 2851 Weather Forecasts (ECMWF) (128, 129). We then used ocean surface currents to calculate eddy kinetic energy (*EKE*) as: $EKE = 0.5 * ((u - \bar{u})^2 + (v - \bar{v})^2)$, where u and v are eastward- and 2852 northward ocean currents respectively and the bar indicates the time-average. All environmental 2853 2854 data were linearly interpolated to 1° (horizontal) resolution.

- 2855
- 2856
- 2857 Model Set

We used the following set of seven models to explain the occurrence of residences and migratory behaviour, each including a different set of the environmental variables we collated (as described in table S8) and specifically avoiding inclusion of correlated variables in the same model:

2861

2862 Model 1: Behaviour ~ sst + u + v + mld + chla + eke + bathymetry + Month

2863 Model 2: Behaviour ~ ssh + u10 + v10 + turbidity + salinity + eke + bathymetry + Month

2864 Model 3: *Behaviour* ~ O2 + vel + vel10 + mld + chla + bathymetry

2865 Model 4: *Behaviour* ~ temp2m + u + v + mld + chla + eke + bathymetry

2866 Model 5: $Behaviour \sim mld + chla + sst$

2867 Model 6: *Behaviour* $\sim u + v + eke + bathymetry$

- 2868 Model 7 (Null model): *Behaviour* ~ 1
- 2869 2870 The response variable "Behaviour" corresponded to grid-cells where residence or migratory behaviour has been detected plus an equal number of grid-cells where presences were available 2871 2872 in our tracking dataset but no behaviour was detected. The total number of grid-cells with 2873 presence and each of the residence or migratory behaviours detected per taxa are shown in Table 2. We compared the predictive ability of models containing different sets of these environmental 2874 2875 variables using the Akaike's information criterion(130). According to the weight of the Akaike's 2876 Information Criteria (wAIC), model 2 was ranked highest for the different behaviours across all 2877 taxa, with the only exception being residency for penguins and turtles (for which the highest ranked models were model 1 and 3, respectively) (table S9). On average, the highest ranked 2878 2879 model for corridors explained 17.8 % of the deviance (ranging from 5.2 % for turtles to 40.1 % 2880 for penguins), while for residences it explained 12 % of the deviance (ranging from 3.1 % for 2881 seals to 24.8 % for polar bears) (Table 2). 2882

2883 Predictions

2884 We used the highest ranked model to predict which grid-cells are likely to be used as residence 2885 or for migration within the entire area where we had occurrence data for each taxon (see 2886 resulting maps in fig. S17). We did this after applying cross validation using a set of 10 (or 5 2887 depending on available data for each taxon) iterations to assess the predictive ability of the 2888 highest ranked models for each taxon. We assessed the predictive ability using the Cohen's 2889 Kappa statistics (K), which measures the agreement between predicted and real (i.e., obtained) 2890 values(131). We then used Landis & Koch (132) criteria to class results into 'no agreement' ($K \leq$ 2891 0), 'slight agreement' ($0 < K \le 0.2$), and at least 'fair agreement' (K > 0.2). Our K values for 2892 corridors averaged at 0.35 (ranging from 0.17 for turtles to 0.56 for penguins) and for residences 2893 averaged at 0.32 (ranging from 0.22 for birds and seals to 0.44 for cetaceans) (Table 2). To 2894 compute the final important marine megafauna areas across all taxa and months to be considered 2895 for the 30% protection (as shown in the right panel of Fig. 3), we used our predictions results 2896 only from models for which K was above 0.2 before applying the optimization algorithm. 2897

2898 **Optimisation algorithm**

2899 To select important marine megafauna areas for protection, we first assigned a score to each 2900 grid-cell based on the detection of key movement behaviours reflecting migratory corridors or 2901 residency areas across taxa. To do this, we first defined Tc and Tr, respectively, as the number of 2902 taxa using the grid-cell as migratory corridor or residence, and then attributed a higher 2903 importance to grid-cells used as residence (i.e., where animals are likely to spend more time)

- when calculating the product between *Tc* and *Tr* to obtain each score per grid-cell, following:
- $Score = 2^{Tc} x 3^{Tr}$

Using this formula, grid-cells receive scores of 2 or 3 if they are, respectively, used as: migratory 2906 2907 corridor or residence by multiple species of only one taxon, and increasingly higher scores if 2908 they are used both as corridors and residencies across multiple taxa (e.g., we obtained a 2909 maximum score of 1944, for grid-cells used by 5 taxa as residence and 3 taxa as corridors) (fig. 2910 S15). We then ordered the grid-cells by descending scores to increasingly select, according to 2911 this ranking, grid-cells currently not (or only partially) protected, until we reached the 30% target 2912 (30% of 71.1 % area covered by our tracking dataset). The resulting selected grid-cells results in 2913 the polygons shown in Figure 3 (and the results from a sensitivity analyses changing the scores 2914 provided to migratory corridors and residences is provided in fig. S16). We repeated this 2915 procedure for the detected movement behaviours based on the probability of each grid-cell to be 2916 used as residence or for migration by each taxon obtained after our modelling procedure 2917 (detailed below). We selected important marine megafauna areas in decreasing order from the 2918 highest probabilities (closest to 1) until the threshold of 30% was reached, and similarly created 2919 the resulting polygons shown in Figure 3 (right panels).

2921 Supplementary Acknowledgements

2922 We thank all who contributed to field work and data collected, specifically including: Antonio P.

Almeida, Alan Aven, Larissa Avens, Jamyle A. F. Batista, Juan A. Bermejo, Soraya C. Bruno,
 Jaqueline C. de Castilhos, Daniel Cejudo, Daniel Devia Cortés, Michael S. Covne, A. Domingo.

Jaqueline C. de Castilhos, Daniel Cejudo, Daniel Devia Cortés, Michael S. Coyne, A. Domingo,
Scott A. Eckert, Mark Fowler, Bruno Giffoni, Kimberly T. Goetz, Kent Hatch, Jefferson Hinke,

2926 Peter Hong, T. Todd Jones, Warren Joyce, Les Kaufman, Nicole Kowalczyk, Eduardo H. S. M.

- 2927 Lima, Pedro López, Anna MacDonnell, Antonio Machado, Maria Ângela Marcovaldi, Rory
- 2928 McAuley, Joanne Braun McNeill, Michael Meyer, P. Miller, Emily R. Nelson, Leif Nøttestad,
- 2929 Fábio L. das C. Oliveira, Nancy Papathanasopoulou, Luis Felipe López-Jurado, Norman
- 2930 Ratcliffe, Juarez T. Scalfoni, Jeffrey Seminoff, Thiago Z. Serafini, Augusto C. C. D. da Silva,
- 2931 Luciano S. Soares, João C. A. Thomé, Wayne Trivelpiece, Marilda I. Weber, Daniel L Webster,
- 2932 Ben G. Weinstein.
- 2933
- 2934 **M. Antonopoulou** acknowledges that the hawksbill satellite tracking project in the Arabian
- region was implemented in cooperation with partner agencies Wildlife and Aquatic Affairs
- 2936 Bureau; in Iran, Department of Environment; in Oman the Ministry of Environment and Climate
- 2937 Affairs and the Environment Society of Oman; in Qatar the Ministry of Environment, Ras Laffan

2938 Industrial City and Qatar University; in the UAE the Environment Agency Abu Dhabi, the

2939 Emirates Marine Environment Group and the Environment & Protected Areas Authority,

2940 Sharjah. Thanks to HE Razan Al Mubarak and Dr. Fred Launay for their support.

2941 G.A. acknowledges LAMAVE staff and volunteers, and would like to thank Mrs Angelique

2942 Songco and the Park Rangers for their collaboration and support in TRNP. Thanks to the Local

2943 Government Units and local communities of Cagayancillo, Talisayan, Malimono, Pintuyan and

2944 San Ricardo. Thanks to Jake Levenson, Steve De Neef and the Pintuyan People's Organization

2945 "KASAKA" who helped with the overall success of the tagging project.

I.A. acknowledges the Basque commercial and recreational fleets cooperated in releasing thetags.

2948 **R.W.B.** acknowledges that the pilot whale tagging was primarily undertaken by Daniel L.

2949 Webster and Gregory S. Schorr. Field work in the Atlantic was undertaken in collaboration with

- 2950 Duke University Marine Laboratory under the supervision of A.J. Read.
- S.E.C. thanks Warren Joyce, Anna MacDonnell, Mark Fowler and Mark Showell for experttechnical assistance.
- G.C. thanks the NSW National Parks and Wildlife Service for their ongoing commitment andlogistical support on Montague Island.
- 2955 A. Chiaradia thanks P. Wasiak, L. Renwick, R. Holmberg, R. Kirkwood, M. Salton, J.P. Robin,

2956 F. Crenner, N. Chatelain, M. Brucker, L. Pelletier, Z. Hogg, T. Shaw, C. McCutcheon, I.

- Zimmer, L. Mandeltort, C. Johnstone, B. Isaac, Environment Protection Agency and MelbourneWater.
- 2959 **R.W.C.** thanks Marcella Harris, the patrollers of DomSeTCO, International Fund for Animal
- 2960 Welfare (IFAW), Georgia Aquarium, and Bureau of Ocean Energy Management (BOEM).

- 2961 E. Cuevas-Flores thanks all Staff, students and volunteers of the Sea Turtle Conservation
- 2962 Program in Pronatura Peninsula de Yucatan, who collaborated in field work for attaching the
- tags. Finally, thanks to the Directions of the Natural Protected Areas CONANP where we
- worked for tagging individuals.
- 2965 S. Diamant thanks Baleines Rand'eau.
- 2966 **K.L.D.** thanks the many organizations and people who contributed to leatherback satellite tag
- 2967 field work: C. Innis, C. Merigo, G. Purmont, M. Leach, A. Myers, M. Dodge, B. Sharp, S.
- 2968 Landry, M. Murphy, G. Tomasian, N. Fragoso, K. Sampson, K. Hirokawa, J. Casey, Rx
- 2969 Smolowitz, J. Casey, S. Leach, J. Wilson, E. Eldredge, V. Saba, T. Sheehan, and the staff of the
- 2970 New England Aquarium Animal Health and Rescue Departments.
- 2971 A.D.M.D. thanks the St Helena National Trust and the Marine Megafauna Foundation.
- 2972 L.L.D. acknowledges Larisa Avens of NOAA Fisheries' Southeast Fisheries Science Center and
- 2973 Kate Mansfield of the University of Central Florida. Thanks to the North Carolina Aquarium at
- 2974 Pine Knoll Shores for collection and rearing assistance and the Sea Turtle Assistance and
- 2975 Rehabilitation (STAR) Center at the North Carolina Aquarium on Roanoke Island for tagging 2976 assistance.
- 2977 **N.E.** acknowledges assistance from staff and volunteers of Statia National Marine Park and St
- 2978 Maarten Marine Park for beach monitoring and satellite tag attachment.
- 2979 C.F. acknowledges field assistance from Christine Larose, Ron & Bozena Summers, A.J. and
- 2980 Camille Lebarbenchon, and M.L.C. donated some geolocators.
- 2981 S.H.F. is grateful to community partners, Levi Qaunaq and Natalino Piugattak from Igloolik, and
- 2982 Noah Ishulutaq and Timeosie Akpalialuk from Pangnirtung, who were responsible for vessel
- 2983 operations. We appreciate the invaluable logistical support provided by the Igloolik and the
- 2984 Pangnirtung Hunters and Trappers Organizations and the Government of Nunavut for cetacean
- 2985 tagging work. We also thank the Canadian Inuit hunters and the Hunters and Trappers
- 2986 Associations and Organizations for assisting with seal capture.
- 2987 M.P.F., C.A.D., and B.F. thank Scott and Sue Tindale, John Annala (MPI), Kina Scollay,
- 2988 Department of Conservation staff, the New Zealand MPI Observer Programme, and other people 2989 who assisted in various ways.
- 2990 S.F. thanks the Secretary-General of the Union of the Comoros Mr. SaidMohamed Ali Said and
- 2991 Direction de l'Environnement, de l'Aménagement et du Logement de Mayotte for issuing the
- scientific permits for this research. Thanks also to the ecovolunteers and eco-tourists for their
- 2993 most valuable help in the field.
- 2994 **C. Garrigue** thanks all the volunteers for their assistance in the field, especially Magaly
- 2995 Chambellant, Dominique Boillon, Claire Bonneville, S. Derville, Ygor Geyer, Rémi Dodemont
 2996 and Véronique Pérard.
- 2997 J.G. and J.M.W. thank the following for assistance in the field: Thomas Robertson, Celine
- 2998 Barre, Mael Imirizaldu, Thomas Vignaud, Tyffen Read, Claude Chauvet, James Hook, Juergen
- 2999 Zier, Greg Skomal, Barry Bruce, Andrew Chin, the Southern and, Northern Province councils of
- 3000 New Caledonia, the Queensland Boating and Fisheries patrol and the Centre de Recherches
- 3001 Insulaires et Observatoire de l'Environment (CRIOBE), and numerous other volunteers. We
- 3002 thank ACREM with the loan of diving equipment and satellite tags from Centre de Recherches
- 3003 Insulaires et Observatoire de l'Environnement (CRIOBE).
- 3004 \

- 3005 K.C.H. thanks Sir Hew Hamilton-Dalrymple and the Scottish Seabird Centre, North Berwick,
- 3006 for access to Bass Rock; and Maggie Sheddan and the Dale family for logistic support.
- 3007 N.H. acknowledges the dedicated contributions of all the University of Miami's Shark Research
 3008 and Conservation Program team members for their assistance.
- 3009 **R.H.** Field support was provided in the Ross Sea by Malcolm O'Toole, Rupert Woods, and
- 3010 Antarctica New Zealand and in Prydz Bay by Malcolm O'Toole, Andrew Doube, Iain Field, and
- 3011 the Australian Antarctic Division. The ARGOS seal tracking and dive data were sourced and are
- 3012 available from the Integrated Marine Observing System (IMOS), NIWA, and LOCEAN. IMOS
- 3013 is a national collaborative research infrastructure, supported by the Australian Government.
- Work was partially funded by an Australian Research Council Linkage Grant to R.H. and David
 Slip (LP110200603).
- 3016 A.R.H. acknowledges Conservation International, WWF-Ecuador, Stanford-TOPP, National
- 3017 Geographic, Galapagos Conservation Trust, Galapagos National Park Directorate, Turtle Island
- 3018 Restoration Network, Galapagos Science Center, Universidad San Francisco de Quito, Blake,
- 3019 Kymberly y George Rapier Charitable Trust, and MigraMar.
- 3020 **N.E. Hussey** would like to thank Peter Darnborough and the fishing crew of the AlleyCat for
- 3021 assistance with catching and tagging sharks.
- 3022 **D.M.P.J.** acknowledges Tre Packard, Mark Healey and the staff and crew from Mikomoto
- 3023 Hammers for their assistance and expertise in the field.
- 3024 A.A.K. would like to thank A Boyd, H Oosthuizen and D Anders from the Department of
- 3025 Environmental Affairs: Oceans and Coasts Branch for permits and operational support; G
- 3026 Oelofse, H Gold and S Liell-cock from the City of Cape Town and S Waries from Shark
- 3027 Spotters, G Cliff, M Dicken and S Dudley from the KwaZulu Natal Sharks Board, M Weisel and
- 3028 W Chivell from the Dyer Island Conservation Trust, the entire crew and support team of the M/V
- 3029 Ocearch and white shark cage diving operators for operational support and assistance in the field.
- 3030 **F.O.L.** acknowledges funding received for the project Reproductive biology, feeding habitats
- 3031 and behavior of the silky shark in the southwestern equatorial Atlantic Ocean. Special thanks
- forwarded to all collaborator's hard work and to those ones that make the work possible at Saint
- 3033 Peter and Saint Paul Archipelago (SPSPA).
- 3034
- 3035 M. Marcoux Narwhal tagging efforts were led by the Department of Fisheries and Oceans long-
- 3036 term marine mammal monitoring program in conjunction with the University of Windsor. All
- field team members of the Ecosystem Approach to Tremblay Sound are thanked for their hard
- 3038 work and commitment in the field especially Bob Hodgson who organized and led the
- 3039 expeditions. We also owe huge gratitude to Jack Orr for his wealth of experience in dealing with
- 3040 narwhal and for leading the expedition equipping narwhal with satellite transmitters during the
- 3041 first research period. Thanks also goes to the Polar Continental Shelf Program and Golder
- 3042 Associates for logistic support in the field.
- 3043 L.M. was principally supported through the Australian Government's Fisheries Research and
- 3044Development Corporation (FRDC) Grants Scheme (PN 2005/031), co-funded by the South
- 3045 Australian Sardine Fishery. We also thank the Nature Foundation South Australia for financial
- 3046 assistance that supported the purchase of GPS units.
- 3047 **G.M.** appreciates the collaboration of the participating fishing vessels.

- 3048 M.M.C.M. acknowledges the logistic support from the Brazilian Navy, the Brazilian Antarctic
- Program (PROANTAR) and SECIRM, the crew from the "Southern Elephant Seal Project"(PEMS) for field support.
- 3051 M.A.C.N. acknowledges the support of numerous staff and volunteers from the Mauritian
- 3052 Wildlife Foundation (MWF) and the National Parks and Conservation Service (NPCS). Of
- 3053 particular note are the contributions made by Vimul Nundlaul, Nicolas Zuel, Richard Baxter, Pat
- 3054 Banville, Katherine Booth Jones, Lucy Rouse and Helen Gath.
- 3055 **B.M.N.** and **S.D.R.** acknowledges the many supporters, funders, donors and volunteers of
- 3056 ECOCEAN Inc. (including the Western Australian Department of Education and all the schools
- 3057 involved in the ECOCEAN Whale Shark Race Around the World), without whom our long-term
- 3058 tagging programme could not have been conducted. We also thank the Western Australian
- 3059 Department of Biodiversity Conservation and Attractions (formerly Department of Environment
- and Conservation and Department of Parks and Wildlife), and all involved in the whale shark
- 3061 ecotourism industry at Ningaloo Marine Park for their ongoing support of our work.
- 3062 E.O. acknowledges the MARECO and Census of Marine Life projects.
- 3063 **S. Oppel** acknowledges the enthusiastic assistance and advice during data collection from
- 3064 Richard Hesketh, Dane Wade, Catherine Supple, Natasha Williams, Kenickie Andrews, Pete
- 3065 Mayhew, and Nathan Fowler. Nigel Butcher and Andrew Asque assisted with preparation of 3066 loggers and equipment, and Elizabeth Marsden kindly provided the base station to download
- 3067 data. We thank G. Balogh, J. Fadely, C. Monnett, J. Gleason, B. Anderson, P. Martin, H. Trefry,
- 3068 S. Trefry, M. Hay, T. Bowman, T. Obritschkewitsch, C. Rea, A. Lazenby, J. Harth, D. Douglas,
- 3069 R. Suydam, D. Troy, J. Zelenak, P. Howey, G. Raven, B. Griffith, and many field assistants for
- 3070 valuable input and technical assistance. E. Taylor initiated the satellite telemetry project, and we
- 3071 appreciate his efforts. L. Phillips assisted in catching birds as well as analyzing telemetry data,
- 3072 and we greatly appreciate her help. We are also grateful to our veterinarians C. Scott, P. Tuomi,
- and M. Mitchell, and several vet technicians for performing the surgeries. Masked booby work
- received enthusiastic assistance and advice during data collection from Kenickie Andrews, Pete
 Mayhew, Phil Lambdon, Shayla Ellick, Dave Higgins, the St Helena National Trust, and staff in
- 3075 Maynew, Phil Lambdon, Shayla Ellick, Dave Higgins, the St Helena National Trust, and sta 3076 the Environmental Management Division of the Environmental and Natural Resources
- 3077 Directorate on St Helena. Nigel Butcher and Andrew Asque assisted with preparation of loggers
- 3078 and equipment. Murphy's petrel work received permission from the Government of the Pitcairn
- 3079 Islands to work on Henderson Island, J. Lavers, L. MacKinnon, A. Forrest, and A. donaldson for
- 3080 assistance in the field, and J. Kelly, A. Bond, J. Vickery, J. Hall, A. Schofield, and C. Stringer
- 3081 for general support. The crew of the Claymore II provided transportation to and from Henderson
- 3082 Island. G. Brodin provided useful support on operation of the GPS devices and data
- 3083 downloading.
- 3084 **A.M.P.** thanks the numerous biologists and pilots who enabled the data collected in this study.
- 3085 The Argos and GPS polar bear tracking data are available through the USGS Alaska Science
- 3086 Center data repository https://doi.org/10.5066/F7RV0MK4. Any use of trade, firm, or product
- names is for descriptive purposes only and does not imply endorsement by the United StatesGovernment.
- **D.M. Parker** acknowledges the US Government through NOAA, and the Governments of Federated States of Micronesia and Republic of Marshall Islands.
- **R.A.P.** thanks all those involved in the long-term tracking and monitoring programmes at Bird Island. John Croxall, Andrew Wood and Janet Silk for overseeing projects, curation, and

- 3093 processing of tracking data. Work represented a contribution to the Ecosystems component of the 2004 DAS Polar Science for Planet Forth Programme funded by NEPC
- 3094BAS Polar Science for Planet Earth Programme, funded by NERC.
- 3095 **M.P.** gratefully acknowledges veterinary assistance from Baukje Lenting and Sarah Michael, and 3096 logistic support from Antarctica New Zealand.
- 3097 D.R. acknowledges field work support by Erick Higuera, Siddharta Velazquez, and Maritza Cruz
 3098 Castillo.
- 3099 R.R.R. and P.J.N.B. thank Dawn Cory-Toussaint, Nadia Hansa, Daniel Kotze, John Dickens,
- 3100 Nasreen Kahn, Rowan Jordaan and Chris Oosthuizen for their efforts in the field. Formerly the
- 3101 Department of Environmental Affairs, now Department of Forestry, Fisheries and the
- 3102 Environment provided logistical support.
- 3103 F.G.R. thanks N. Veiga, J. L. Veiga, G.Pereyra, L. Olivera, D.J. Shuman, H. Katz, R. Frau, S.
- 3104 Tavoni, M. Garcia, M. Rivas for their invaluable assistance and collaboration during the field
- 3105 work. We are also thankful to C. Leiza (Parque Zoologico Lecoq, Uruguay) for providing
- 3106 tranquilizing darts, and the Marine Mammal Center (Sausalito, California, USA) for lending the
- 3107 capture nets.
- 3108 **Y.R.** thanks several students and collaborators for their contribution to the data collection.
- 3109 **K.S.** thanks the Department of Environment, Forests and Climate Change, Andaman, and
- 3110 Nicobar Islands for their logistical support. We are deeply grateful to all the local field staff who
- 3111 accompanied us for our monitoring camps and surveys.
- 3112 G.L.S. thanks the Leatherback Trust and the Betz Chair of Environmental Science at Drexel
- 3113 University for their assistance. The Costa Rican Ministry of Natural Resources and the
- 3114 Environment provided research permits. We thank G. Goldring, the Goldring Marine Biology
- 3115 Station, and the staff and volunteers at PNMB for support at Playa Grande.
- 3116 L.L.S. thanks G. Hays, J. Houghton and T. Doyle for tagging two sunfish in Ireland; and all the
- 3117 crew in Tunipex S.A., especially both Mr Morikawa and captain Alfredo Poço, for allowing
- 3118 access to the tuna pen and for much valued field support.
- **J.T.S.** acknowledges NOAA, University of California Santa Cruz, Dr. Sara Iverson, Alison
- 3120 Banks, and Tonya Zeppelin.
- 3121 A. Takahashi thanks the Native Village of Savoonga for approving fieldwork, and Michael
- 3122 Toolie, Punguk Shoogukwruk, M. Ito, Y. Suzuki, Y. Watanuki, T. Yamamoto, N. Oka, N.
- 3123 Katsumata, K. Sato., S. Watanabe, K. Watanabe, Y. Hirose, T. Fukuda, S. Sasaki, K. Tamori, D.
- 3124 Ochi, and S. Hirose for their assistance with fieldwork.
- 3125 P.M.T. acknowledges the Orkney Islands Council and National Trust for Scotland for access to3126 field sites.
- 3127 M.T. acknowledges Woodside Energy Ltd (Woodside) as Operator for and on behalf of the
- 3128 Greater Enfield Project with Mitsui E & P Australia Pty Ltd and the Browse Joint Venture.
- 3129
- 3130 **P.N.T.** thanks all those associated with the long-term monitoring and tracking programmes at
- 3131 Bird Island and Signy, including those responsible for data curation and validation. Data used
- 3132 here were a contribution to the BAS Ecosystem programme, funded by UKRI/NERC.
- 3133
- 3134 N.V. acknowledges Project "Life B4-3200/97/247 to support the conservation of the bottlenose
- dolphin and the loggerhead turtle in the Canary Islands"/"Life B4-3200/97/247 de apoyo a la

- 3136 conservación del delfín mular (Tursiops truncatus) y la tortuga común (Caretta caretta) en
- 3137 Canarias" (tracking period 1998-2000), Project "Aegina (Interreg IIIB 04/MAC/3.5/C36)"
- 3138 (tracking period 2006-2009), and Monitoring plan of the loggerhead turtle in the Canary Islands"
- 3139 / "Plan de seguimiento de la tortuga boba en Canarias" (tracking period 2008-2010; plan
- 3140 executed by the Grandilla Environmental Observatory OAG, Observatorio Ambiental
- 3141 Granadilla), plus all the collaborative work with the University of Las Palmas de Gran Canaria
- and the Spanish Herpetological Society as well as with the Instituto Canario de Ciencias Marinas
- 3143 (ICCM) of the Gobierno de Canarias, the Instituto Nacional de Desenvolvimento das Pescas
- (INDP), and the Direcção Geral do Ambiente, the latter depending on the Ministerio de
 Ambiente, Agricultura e Pescas of the Republic of Cabo Verde. Thanks to the wildlife recovery
- 3145 Ambiente, Agricultura e Pescas of the Republic of Cabo Verde. Thanks to the wildlife recovery 3146 center of Tafira, Cabildo de Gran Canaria, and the ONG Cabo Verde Natura 2000 as well as,
- 3147 Cabildo de Fuerteventura; and the Society for the Study of Cetacean in the Canary Archipelago
- 3148 (SECAC, Sociedad para el Estudio de los Cetáceos en el Archipiélago Canario) for logistic
- 3149 support "). N.V. also thanks Daniel Cejudo, Luis Felipe López-Jurado, Juan A. Bermejo, and
- 3150 Antonio Machado for assistance in the field.
- 3151 S.V. thanks The Charles Darwin Foundation, Charles Darwin Research Station, and Parque
- 3152 Nacional Galapagos for their logistics and fieldwork support. We also thank P. Howorth and the
- 3153 volunteers from the Marine Mammal Center in Santa Barbara; E. Stetson, A. Parás, B.
- 3154 McDonald, D. Páez, C. Martínez and volunteers from the Darwin Station; J. Torres, D. Schuman,
- 3155 S. Barberán, D. Aurioles, F. Trillmich, S. Salazar, D. Casper, M. Zavanelli, L. Huckstadt, C.
- 3156 Kuhn, V. Michuy, M. Rutishauser, P. Robinson, S. Simmons, and the San Cristóbal naval base
- 3157 for their help in the field.
- 3158 C.V. acknowledges field work involved many participants belonging to the Sea Mammal
- 3159 Research Unit, University of La Rochelle (LIENSs, PELAGIS, CEBC), Aerobaie, Réserve
- 3160 Naturelle Domaine de Beauguillot, Direction régionale de l'environnement, de l'aménagement et
- 3161 du logement (DREAL) Basse Normandie, Office National de la Chasse et de la Faune Sauvage
- 3162 (ONCFS), Zoo de la Flèche, Zoo de Vincennes, Zoo la Bourbansais, Picardie Nature, Parc
- 3163 naturel marin d'Iroise, Oceanopolis, and Université de Lièges (Belgium).
- 3164 S. Wanless acknowledges field assistance provided by Stuart Murray and Jill Harden. Sirtrack,
- seaturtle.com, and Ewan Wakefield provided advice and support for logger deployments andpost-processing.
- 3167 **N.E.W.** acknowledges that tags were included in her PhD thesis at James Cook University,
- 3168 supported by the JCU Postgraduate Research Scholarship.
- 3169
- 3170 L.J.W. acknowledges that northern Gannet data were collected in a study led by Rowena H.W.
- 3171 Langston with Emma Teuten leading the data management and processing. John Hartley
- 3172 managed the project funding on behalf of DECC and Philip Bloor supported DECC's funding of
- 3173 the work. Work was made possible thanks to collaboration between the Joint Services Mountain
- 3174 Training Wing, Defence Training Estate North, the University of Leeds, East Yorkshire Ringing
- 3175 Group, RSPB staff and volunteers. We are particularly grateful to Captain R. Groves RAPTC,
- 3176 Captain S Higgins APTC, QMSI Haslam, Andy Phillips, Major Tony Crease (Retired), Tony
- 3177 Haw, Paul "Chips" Rafferty, Professor Keith Hamer, David Aitken, John Bell, Chris Blakeley,
- 3178 Sergio Boggio, M.B., Chris Bradshaw, Paul Britten, Nigel Butcher, Keith Clarkson, Ian Dillon,
- 3179 Peter Dunn, Chris Hansell, Ian Kendall, Reg Langston, John McEachen, Linda McKenzie, Lucy

- 3180 Murgatroyd, Joanne Peyton, Ruth Porter, Steve Race, Sophie Rainer, Zoe Tapping, Mark
- 3181 Thomas, Paul Thorpe and Ewan Wakefield.
- 3182
- 3183 **D.J.Y.** is thankful for support from the Resolute Bay Hunters and Trappers Association and
- 3184 community partners, especially Peter, Jeff and Uluriak Amarualik.
- 3185 P.M.Z. acknowledges field assistance by L. Díaz, D. Fuenzalida, E. Garcés, P. Ojeda, L. Pizarro
- 3186 and the crew of the Vama II.
- 3187

3188 Supplementary Author Contributions

- 3189 Contact Author for Access to Tracking Data
- 3190 S.A., F.J.A., A.S.A., P.A., S.Å., J.A.A., G.A., J.M.A., I.A., R.W.B., G.H.B., A. Barnett, A.
- 3191 Beard, E.J. Belda, S.R.B., D.B., M.L. Berumen, S. Bessudo, B.A.B., R.B., C.D.B., J.M.B.,
- 3192 S.E.C., H.A.C., A. Carlisle, R.H.C., G.C., P.C., L.C.F., D.D.C., A. Chiaradia, C.R.C., C.C.,
- 3193 E.E.G.C., R.C., D.P.C., M.C., E. Cuevas-Flores, L.D., R.D., P.J.N.B., T.D., K.L.D., T.K.D.,
- 3194 C.M.D., L.L.D., N.E., S.H.F., B.F., R.J.F., S.F., A.S.F., C. Garrigue, E.G., B.J.G., S.D.G.,
- 3195 V.G.C., W.J.G., J.R.G., T.L.G., H.M.G., N.H., M.O.H., R.H., C.E.H., A.R.H., R.W.H.I., B.J.H.,
- 3196 L.A. Howey, R.E.H., N.E. Hussey, P.H.L., C.H., D.M.P.J., A.J., M.J., A.A.K., F.O.L., M.L.C.,
- O.A.L., J.J.L., G.L., M. Lopez Mendilaharsu, P.L., M.E.L., B.C.L.M., A.I.M., M.L. Mallory,
 J.C.M., D.M., M. Marcoux, B.M., D.A., R.L.M., L.M., J. Morris, M.A.C.N., B.M.N., E.O., S.
- 3199 Oppel, A.M.P., B.P., V.H.P., D.M. Palacios, C.P., L.R.P., C.R.P., R.A.P., N.J.P., A.N.P., R.P.,
- 3200 N.Q., J.L.Q., I.R., D.R., J.A.R., A.F.R., R.R.R., F.G.R., D.P.R., T.L.R., C.A.R., Y.R., D.R.L.R.,
- 3201 E.A.P.S., G. Schofield, E.S., S.A.S., K.S., G.L.S., M.S.S., M.A. Silva, D.W.S., G. Skomal, M.
- 3202 Soria, J. Stahl, J.D. Stewart, A. Takahashi, V.T., P.M.T., M.T., J. Tomás, P.N.T., F.V., N.V.,
- 3203 S.V., C.V., S. Wanless, M.J. Weise, R.S.W., S.D.W., N.E.W., D.N.W., S. Wischnewski, M.J.
- 3204 Witt, D.J.Y., P.M.Z., A.Z.
- 3205
- 3206 Led, Designed or Secured Funds for Fieldwork
- 3207 F.J.A., A.S.A., P.A., S.Å., R.D.A., M. Antonopoulou, J.A.A., G.A., J.M.A., H. Arrizabalaga, S.
- 3208 Bach, R.W.B., G.H.B., S.G.B., A. Barnett, W.B., A.M.M.B., A. Beard, E.J. Belda, I.B., A.
- 3209 Bennison, S.R.B., D.B., M.L. Berumen, S. Bessudo, N.P.A.B., M.E. Bond, R.B., C.D.B., A.C.B.,
- 3210 M.L. Brooke, E.J. Brooks, J.M.B., H.A.C., R.A.C., A. Carlisle, R.H.C., G.C., P.C., F.R.C.,
- 3211 L.C.F., D.D.C., T.K.C., J. Charrassin, A. Chiaradia, C.R.C., C.C., E.E.G.C., J.E.M.C., R.C.,
- 3212 D.P.C., M.C., E. Cuevas-Flores, K.P.D., R.D., P.J.N.B., K.L.D., M.C.D., A.D.M.D., T.K.D.,
- 3213 M.J.D., L.L.D., C.A.D., P.H.D., E.W.J.E., L.D.E., M.V.E., N.E., A.I.F., C.F., S.H.F., J.
- 3214 Filmalter, G.C.F., R.J.F., A.F., S.F., M.P.F., A.S.F., A.J.G., C. Garrigue, E.G., H.G.G., B.J.G.,
- 3215 S.D.G., V.G.C., W.J.G., C. Guinet, J.G., T.L.G., H.M.G., K.C.H., N.H., R.H., E.H., C.E.H.,
- 3216 G.H., G.C.H., M.H., A.R.H., M.P.H., R.W.H.I., A.E. Herrera, M.A.H., B.J.H., L.A. Howey, L.A.
- 3217 Huckstadt, R.E.H., N.E. Hussey, C.H., D.M.P.J., A.J., M.Y.J., M.J., O.J.D.J., R.J., S.J.J., A.A.K.,
- 3218 P.K., F.O.L., M.L.C., J.L., J.J.L., G.L., G.G.L., M. Lopez Mendilaharsu, A.D.L., M.E.L.,
- 3219 W.S.L., B.C.L.M., A.I.M., C.A.M.H., M.L. Mallory, A.M., M. Marcoux, D.A., H. Marsh,
- 3220 J.D.M., R.L.M., L.M., C.R.M., M.G.M., G.M., M.M.C.M., W.J.N., M.A.C.N., B.M.N., K.N.,
- 3221 L.N., S. Oppel, B.P., V.H.P., L.R.P., C.R.P., J.G.P., R.A.P., S.J.P., N.J.P., P.P., M.P., A.N.P.,
- 3222 T.J.P., R.P., N.Q., J.L.Q., T.R., D.R., R.R., A.F.R., R.D.R., R.R.R., F.G.R., D.P.R., P.W.R.,
- 3223 T.L.R., C.A.R., Y.R., P.M.S., S.S., G. Schofield, J.M.S., E.S., S.A.S., K.S., G.L.S., M.S.S., M.A.
- 3224 Silva, D.W.S., G. Skomal, D.J.S., L.L.S., J. Stahl, J.T.S., G.M.W.S., J.D. Stewart, A.
- 3225 Swaminathan, A. Takahashi, V.T., P.M.T., S.R.T., M.T., J. Tomás, P.N.T., R.S.B., N.V., M.
- 3226 Vely, S.V., C.V., S. Wanless, S.B.W., N.W., M.J. Weise, L.W., R.S.W., B.M.W., S.D.W., B.W.,
- 3227 D.N.W., A. Will, S. Wischnewski, M.J. Witt, J.C.X., T.Y., D.J.Y., P.M.Z., A.Z., A.N.Z.
- 3228
- 3229 Undertook Animal Tagging
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3327 Fig. S1. Total number of tracked individuals per latitude and longitude

3328 Spatial extent of the marine megafauna tracking datasets including the 12,794 individuals in the

3329 global dataset at 1° resolution, with top and left inset plots representing longitudinal and

latitudinal coverage of the curated tracks with dotted histograms showing the number of

individuals tagged (at tagging locations) and shaded areas indicating the number of individuals

3332 with tracked positions in the same geographical locations. These plots show that a higher number

- 3333 of tracked individuals is not necessarily related to tagging locations.
- 3334
- 3335



Fig. S2. Biodiversity captured versus expected based on the 111 species considered in our dataset.

3339 Map shows that our dataset captures some of the known hotspots, including in the NE Pacific,

3340 NW Atlantic, and regions of the SW Atlantic, but it misses others (e.g., Indo-Pacific and Central

West Pacific regions, Gulf of Guinea, and waters around Australia and Madagascar). A) Global map depicting the first locations tracked per individual, providing a visual representation of the

tagging locations. B) Effective number of species (S_{eff}) observed in each grid-cell (*c*) where we

- had tracking data based on the Shannon entropy (H)(31). H was calculated from the probability
- 3345 of observing each species among all the individuals visiting each grid-cell $(p_s(c))$, which in turn
- 3346 is the result of the fraction of tracked individuals $f_s(c)$ of each species (s) grid-cell with at least

3347 one location within each 1° grid-cell divided by the total number of tagged individuals of the

3348 same species. The resulting fraction is independent from the tagging effort excluding potential

biases arising from the different number of tagged individuals of each species.

3350
$$S_{eff}(c) = 2^{H(c)}$$
 where $H(c) = \sum_{s} p_{s}(c) \log_{2} p_{s}(c)$ and $p_{s}(c) = \frac{f_{s}(c)}{\sum_{s} f_{s}(c)}$

3351 C) Expected richness hotspots for the species considered based on species geographical range

3352 shapefiles obtained from the iucnredlist.org/ (accessed 24 Jan 2022) for all species, except for

3353 flatback turtles, which were obtained from the Recovery Plan for Marine Turtles in Australia

(2017)(133). For comparison with global biodiversity maps see literature for birds(134),

mammals(51), sharks(14), and also general marine biodiversity (i.e., also including plants, corals
and non-migratory animals)(135).



Fig. S3. Marine Protected Areas (MPAs) and Exclusive Economic Zones (EEZs) in the marine environment

- We obtained geographical information defining existing marine protected areas (MPAs,
- including marine parks; shown in blue) from protectedplanet.net (accessed 10 June 2021) (33)
- and exclusive economic zones (EEZs; shown in grey) from marineregions.org/ (accessed 28 June 2021) (34).





3369 Fig. S4. Number of species affected by each anthropogenic threat considered

3370 Summaries of the number of species known to be impacted by each threat considered in our 3371 analyses based on the listed threats and sub-threats on the IUCN Threats Classification Scheme 3372 v3.3(50) as detailed in table S4. Figure shows commercial fishing and climate change 3373 (represented as SST anomaly) has having impacts on the highest number of species analysed in 3374 this study, and with fixed gear and longlines fishing gears listed for most species. All sea turtles, 3375 sirenians and polar bears are affected by most of the threats. A large number of birds, cetaceans 3376 and fishes are impacted by fishing and SST anomaly, with higher number of birds and fish 3377 species being impacted by plastic, and higher number of cetacean species being the most impacted by shipping and noise. Species considered in each taxon group include flying birds 3378 3379 (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, 3380 polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and manatees), and turtles.



3381 Fig. S5. Assessment of behaviours detected when using only one position per day for each individual

3382 Results for the final tracking dataset when considering only one position per day per individual across all taxa. All maps show similar 3383 spatial patterns to those obtained for the complete dataset (as shown in Fig. 1 and Fig. S2). A) Top: Total number of individuals for 3384 which we have tracking data in each grid-cell; Bottom: effective number of species obtained per grid-cell. B) Top: Identified 3385 migratory and residence areas when using only one position per day per individual across all taxa (greyed grid-cells indicate that no 3386 key movement behaviour was identified); Bottom: Results obtained when using the Jaccard Index to compare the results obtained here 3387 with those obtained from the original dataset. Species considered in each taxon group include flying birds (listed as birds), cetaceans 3388 (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and 3389 manatees), and turtles.



3391 Fig. S6. Assessment of behaviours detected when removing the tagging location bias

Results for the final tracking dataset after removing the tagging location bias. All maps show similar spatial patterns to those obtained for the complete dataset (as shown in Fig. 1 and fig. S2). A) Top: total number of individuals for which we have tracking data in each grid-cell; Bottom: effective number of species obtained per grid-cell. B) Top: Identified migratory and residence areas when removing all locations around the tagging location for all individuals across all taxa (greyed grid-cells indicate that no key movement behaviour was identified); Bottom: Results obtained when using the Jaccard Index to compare the results obtained here with those obtained from the original dataset. Species considered in each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (*Ursus maritimus*), seals, sirenians (i.e., dugongs and manatees), and turtles.



3400 Fig. S7. Assessment of behaviours detected when changing resolution to 0.5°

Results for the final tracking dataset when considering a spatial resolution of 0.5° All maps show similar spatial patterns to those obtained for the complete dataset (as shown in Fig. 1 and fig. S2). A) Top: total number of individuals for which we have tracking data in each grid-cell; Bottom: effective number of species obtained per grid-cell. B) Top: Identified migratory and residence areas when halving the spatial resolution (greyed grid-cells indicate that no key movement behaviour was identified); Bottom: Results obtained when using the Jaccard Index to compare the results obtained here with those obtained from the original dataset. Species considered in each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (*Ursus maritimus*), seals, sirenians (i.e., dugongs and manatees), and turtles.



Fig. S8. Assessment of ecological meaningful behaviour when changing resolution to 2° 3409

3410 Results for the final tracking dataset when considering a spatial resolution of 2°. All maps show similar spatial patterns to those obtained for the complete dataset (as shown in Fig. 1 and fig. S2). A) Top: total number of individuals for which we have tracking 3411 3412 data in each grid-cell; Bottom: effective number of species obtained per grid-cell. B) Top: Identified migratory and residence areas 3413 when doubling the spatial resolution (greyed grid-cells indicate that no key movement behaviour was identified); Bottom: Results 3414 obtained when using the Jaccard Index to compare the results obtained here with those obtained from the original dataset. Species 3415

considered in each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly

3416 sharks), penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and manatees), and turtles.

Birds



Fishes



Polar bears





Penguins



Seals



Sirenians



Turtles





3417 Fig. S9. Detected important migratory corridors and residence areas found by taxa

3418 Spatial representation of migratory corridors and residence areas by taxa detected based on our 3419 analyses of coherence and z-scores (Methods). Results include the migratory corridors shown

- with faded colours for each taxon, which were obtained after detection of the time window that
- with faded colours for each taxon, which were obtained after detection of the time window that

- resulted in the maximum number of 1° grid-cells showing coherent direction of movement 3421
- 3422 within each month for each taxon (fig. S12), and after detection of hotspots of coherence clusters
- using on a Lorenz curve approach (fig. S13). Residence areas, indicated by grid-cells shown in 3423
- 3424 stronger colours, were obtained based on z-scores of one standard deviation below the average
- displacement for each taxon. Where both migratory corridors and residence areas were found, 3425
- 3426 grid cells include a black outline. Grey indicates grid-cells where no specific behaviour was
- 3427 identified. White areas depict regions without tracking data. Species considered in each taxon
- 3428 group include flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and
- 3429
- 3430 manatees), and turtles.



3432 Fig. S10. Statistics of the space and time used for different behaviours per taxa

3433 Fractions of space use (left panel) and time spent (right panel) in different behaviours calculated

3434 for each individual to show the distribution of the results obtained per taxon. Shown are the

3435 median values (percentile 50, dots) connecting the percentiles 25 and 75 obtained across

3436 individuals within each taxa. These values represent the most likely values on any sample of

3437 tracked individuals (with other values obtained outside the interval between percentiles 25 and

3438 75). Species considered in each taxon group include flying birds (listed as birds), cetaceans

3439 (mostly whales but also dolphins), fishes (mostly sharks), penguins, polar bears (Ursus

3440 *maritimus*), seals, sirenians (i.e., dugongs and manatees), and turtles.



Fig. S11. Global footprint of anthropogenic threats and intensity within important areas for marine megafauna (IMMegAs)

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The global footprint of anthropogenic threats is displayed as global averages per threat. Threat intensity outside the important marine megafauna areas identified in this study (left: based on the movement data, and right: based on model predictions) is indicated by the grey colour bar, and by the coloured bars per threat within important areas. Threats depicted include, from top to

- bottom: fishing intensity, shipping density, plastic density, and warming (according to anomalies
- 3451 to sea surface temperature; SST).
- 3452





3455 Fig. S12. Average monthly coherence results at multiple time windows for all taxa

3456 Our analyses for detection of key movement behaviours indicated by migratory or residency-like 3457 behaviours showed that the maximum number of 1° grid-cells with coherent movement was 3458 obtained for time windows (Tw) < 10 days for all taxa considered: 2 days for fish, 3 days for 3459 birds, 4 days for cetaceans, 6 days for penguins, polar bears, seals, and turtles, and 7 days for 3460 sirenians. Circles indicate the time at which coherence was highest for each taxon. Species 3461 considered in each taxon group include flying birds (listed as birds), cetaceans (mostly whales 3462 but also dolphins), fishes (mostly sharks), penguins, polar bears (Ursus maritimus), seals, 3463 sirenians (i.e., dugongs and manatees), and turtles.





3465 Fig. S13. Detection of cluster hotspots of coherence with the Lorenz curve

3466 Plots show the results obtained per taxon, when considering multiple temporal scales of 1, 2, 3, 4, 5, 6, and 12 months to identify hotspots of clusters of grid-cells with coherent movement (i.e., 3467 grid-cells for which the cosine of the angle between their average directions is > 0.8) that 3468 3469 resulted in the global migratory corridors shown in Figure 3 (also shown per taxon in Fig. S9). 3470 The threshold for minimum cluster size defining a migratory corridor at each temporal scale for 3471 each taxon was identified with a Lorenz curve (parameter free approach; 121). For example, for 3472 birds, the hotspots for minimum cluster size were detected at the top 0.5% of the cumulative 3473 distribution (i.e., for $x_{th} = 0.995$), indicating the minimum cluster size to identify a migratory 3474 corridor was 52 connected grid-cells. Migratory corridors were therefore, defined at minimum 3475 cluster sizes (S_{th}) of 52 grid-cells for birds, 23 for cetaceans, 17 for fishes, 8 for penguins, 5 for polar bears, 40 for seals, and 29 grid-cells for turtles. No hotspots of clusters of grid-cells with 3476 coherent movement were detected for sirenians, likely due to lack of data. Species considered in 3477 3478 each taxon group include flying birds (listed as birds), cetaceans (mostly whales but also 3479 dolphins), fishes (mostly sharks), penguins, polar bears (Ursus maritimus), seals, sirenians (i.e.,

3480 dugongs and manatees), and turtles.



3482 Fig. S14. Comparison of detected residency areas across taxa after randomising tracks

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3483 Maps show the grid-cells identified as residency areas for each taxon after (a) and before (b) one 3484 example of randomised tracks (see Materials and Methods) to demonstrate animals are 3485 selectively using space. Patterns shown on the bottom map follow those identified in the original 3486 dataset and displayed in Fig. 2. (c) shows the Jaccard index (i.e., area of overlap divided by area 3487 of union) obtained for each taxon. The low values for the index across taxa confirm the 3488 independence of the results before and after randomising the dataset. Colours refer to birds (light 3489 green), cetaceans (dark yellow), fishes (red), penguins (dark green), polar bears (orange), seals 3490 (blue), sirenians (purple), and turtles (pink). Species considered in each taxon group include 3491 flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), 3492 penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and manatees), and 3493 turtles.



Fig. S15. Depiction of scores computed as part of the optimisation algorithm to select 3495

3496 priority areas for global protection of marine megafauna

3497 Areas used by multiple taxa (noting that each taxa group includes multiple marine megafauna

3498 species, table S1) are indicated with warmer colours (i.e., in purple, orange and red according to

3499 an increase in taxa and behaviours observed).





- 3503 Results obtained when the scores for migratory corridors (S_C) and for residences (S_R) used to run the optimization algorithm were
- 3504 changed: (Top left) $S_R=3$ and $S_C=2$ as used throughout the manuscript, (Top rigth) $S_R=2$ and $S_C=2$, (Bottom left) $S_R=2$ with no score
- 3505 for corridors, and (Bottom right) $S_C=2$ with no score for residence areas (and showing the most different results as expected).




























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Fig S17. Comparison of detected and predicted areas used for important marine megafauna behaviours

3522 Shown are the maps we detected by our direct analyses of the tracking data (left) and those based on predicted probabilities of behaviours occurring (right) for residency (top) and migratory 3523 behaviours (bottom) for each taxon. Asterisks are included in the predicted maps for seal 3524 3525 residences and turtle corridors because the models leading to these predictions results in a K < 0.23526 and predictions were therefore, not considered when merging important marine megafauna areas 3527 across all taxa based on modelled probabilities. Species considered in each taxon group include 3528 flying birds (listed as birds), cetaceans (mostly whales but also dolphins), fishes (mostly sharks), 3529 penguins, polar bears (Ursus maritimus), seals, sirenians (i.e., dugongs and manatees), and 3530 turtles.