Title: Mitochondrial phylogeography reveals high diversity and unique divergent lineage in Indian Dugongs (*Dugong dugon*)

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ABSTRACT:

- 1) India plays a central role in dugong conservation by hosting the largest population within south Asia. Current knowledge on status of Indian dugongs is limited due to paucity of reliable ecological data. This study generates mitochondrial control region sequences from about 10% of dugong population from major dugong populations within India. These data was compared with the global data to assess genetic lineages, population structure and genetic diversity of Indian populations.
- 2) Multiple analyses suggest that the Indian dugong populations are part of a single genetic cluster, comprising south Asia, northwest Indian ocean and southwest Indian ocean populations. Despite small population size, they retain high genetic diversity with unique mitochondrial DNA haplotypes within south Asia.
- 3) Within India, novel haplotypes were observed from all sampling sites with overall high haplotype diversity (0.85±0.04) but low nucleotide diversity (0.005±0.001). Indian populations exhibit high genetic differentiation with higher within-population variance (63.41%) than among populations (36.59%), signaling population structure. Few haplotypes were shared with Sri Lanka and southeast Asian populations, indicating potential genetic connectivity.
- 4) Being the most genetically unique population within south Asia, Indian dugong populations are globally significant. We recommend that Indian Dugong populations should be managed as a Conservation Unit to ensure population recovery and long-term survival of the species.

KEYWORDS:

Sirenians; genetic lineage, demography, population structure, genetic diversity; marine mammal conservation

1. INTRODUCTION

Dugong (Dugong dugon (Müller)) or sea cow is the only strictly marine herbivore species of the order Sirenia. Historically, dugongs were distributed across the tropical and subtropical regions of the Indo-Pacific Ocean, inhabiting shallow coastal waters ranging from east coast of Africa to western Pacific Ocean (Marsh, Penrose, Eros & Hugues, 2002). Currently classified as 'vulnerable' by the IUCN Red List, their distribution is restricted to Australia, parts of southeast Asia, Indian subcontinent, Arabian Gulf and eastern coast of Africa (Marsh & Sobtzick, 2015). Within their range, Australian coast retains the largest dugong population, followed by Arabian Gulf. All remaining dugong populations across southeast Asia, Indian sub-continent and east African coast are small and fragmented (Hines et al., 2012; Marsh & Sobtzick, 2015). Their coastal distribution, exclusive dependence on seagrass habitats, long lifespan and slow reproduction rates make them vulnerable to human-mediated impacts across their range. Major threats for dugongs include accidental entanglement in fishing nets, hunting/poaching for meat, vessel strikes, degradation of seagrass habitats and coastal infrastructure development (Marsh, O'Shea & Reynolds III, 2011; Hines et al., 2012; Sivakumar, 2013; D'Souza, Patankar, Arthur, Alcoverro & Kelkar, 2013). Despite various protection measures implemented internationally, their numbers are declining across their range with local extinctions from Mauritius, Maldives and Taiwan (Marsh & Sobtzick, 2015).

India retains the largest dugong population in the south Asia sub-region and thus plays a significant role in dugong conservation at regional and global scale (Sivakumar, 2013). Recent estimates indicate less than 200 individuals distributed in isolated populations along Gulf of Kachchh (Gujarat, west coast), Gulf of Mannar & Palk Bay (Tamil Nadu, south-east coast) and island archipelago of Andaman & Nicobar (Pandey, Tatu & Anand, 2010; Sivakumar, 2013;

Sivakumar & Nair, 2013; D'Souza, Patankar, Arthur, Alcoverro & Kelkar 2013). With a decreasing population trend and regionally 'Endangered' status, it is absolutely critical to focus on reviving their population and restoring seagrass habitats. However, current ecological knowledge on status of Indian dugongs is limited due to low population densities, rare sighting records and inadequate spatial survey efforts. Lack of long-term monitoring and systematic sampling has also resulted in paucity of reliable information on their genetic lineage in comparison to other dugong populations. So far, genetic status of dugong populations is known from published studies in Australia (Tikel, 1997; McDonald, 1997; Seddon et al., 2014, Blair et al., 2014), Thailand (Palmer, 2004; Bushell, 2013) and western Indian ocean (Plon, Thakur, Parr & Lavery, 2019). These studies indicate three major dugong genetic lineages viz. Australian (restricted and widespread), south-east Asian and western Indian Ocean (Blair, Marsh & Jones, 2013; Plon, Thakur, Parr & Lavery, 2019). Recent work by Plon, Thakur, Parr & Lavery (2019) suggests that the Sri Lankan dugongs are genetically divergent from other populations, based on very limited samples (n=4). Given rapidly declining populations of dugongs in south Asia with recent local extinctions in Mauritius, Maldives and parts of Indian and Sri Lankan coasts (Marsh, Penrose, Eros & Hugues, 2002), adequate sampling to ascertain their position vis-à-vis global populations is of critical importance.

In this paper, we conducted comprehensive genetic sampling of Indian dugongs to describe a) genetic lineage and phylogeography of the Indian dugongs in relation to other dugong populations at global scale and b) the genetic diversity, differentiation and demographic patterns of dugong populations in India. We believe that the results from this study will be critical in developing population specific management plans and thus help in long-term conservation of this globally threatened marine mammal.

1. MATERIALS AND METHODS

2.1 Permits

The permission to carry out fieldwork and genetic sampling of dugongs was provided by the Ministry of Environment, Forest and Climate Change, Government of India (CAMPA Authority letter number: 13-28(01)/2015-CAMPA). Due to non-destructive sampling approach used in this study, no ethical committee approvals were needed.

2.2 Sample collection

Given low population size of dugongs in the Indian subcontinent (<200 individuals), fragmented distribution along the coastal region and rare live sightings (Sivakumar, 2013), it was logistically difficult to conduct systematic sampling of biological material. Hence, most of the sampling was opportunistically conducted from dead stranded dugongs from coastal areas of Gulf of Mannar, Palk Bay, Gulf of Kachchh and Andaman & Nicobar Islands. All samples were stored in ethanol at the field sites and later shipped to Wildlife Institute of India for storage at -20°C until further analysis. In addition to opportunistic sampling, historical samples (bone scrapings) were collected with associated geo-location information from State Forest Departments' collections. The details of the samples and their geographical sampling location are provided in Supplementary Table 1.

2.3 DNA extraction, marker selection and PCR amplification

Total genomic DNA was isolated from all fresh tissue samples using standard protocols mentioned in DNeasy blood and tissue kit (Qiagen, Germany). However, for poor quality museum samples, a modified protocol was used to extract DNA (Mondol, Brufford & RamaKrishnan, 2013). In brief, all samples were cut into small pieces, washed with EDTA and

macerated into tiny fragments. The fragments were then completely digested with 40 µl proteinase K for two days, followed by DNA extraction using DNeasy blood and tissue kit (Qiagen, Germany). For museum samples, extractions were carried out in a space dedicated to low quality DNA samples where no earlier dugong DNA extraction has been conducted. Negative controls were included for every batch of extractions to monitor any possible contamination.

We used a universal mammalian primer (A24- Kocher et al., 1989) and dugong-specific primers A58, A77 and A80 (Tikel, 1997) to amplify the mitochondrial DNA (mtDNA) control region from dugong samples. Post-temperature standardizations of these primers, PCR reactions were performed in 10 μl volume with 5 μl Qiagen multiplex PCR mixture (Qiagen, Germany), 0.2 mg/ml Bovine Serum Albumin (BSA), 0.5μM primer mixture and 2μl (2-40 ng/μl concentration) of DNA. PCR conditions included an initial denaturation of 96°C for 15 minutes, followed by 35 cycles of denaturation (96°C for 30 s), annealing (45°C for 30 s) and extension (72°C for 60 s); followed by final extension (72°C for 10 min). During all amplifications, PCR positive and negative controls were included to monitor any possible contamination.

However, it was challenging to amplify the above-mentioned markers in poor quality/degraded bone samples collected in the study. We designed primers to amplify small amplicon sizes from dugong DNA. All published whole mtDNA sequences were downloaded from GenBank with Accession numbers AY075116.1, AJ421723.1, NC003314.1 and were aligned with the sequences generated from tissue samples collected in this study using MEGA v.6.0 (Tamura, Stecher, Peterson, Filipski & Kumar, 2013). Conserved regions within the sequences were visually identified and primers were designed to amplify <250 bp amplicon size to ensure high amplification success from degraded samples. Post-temperature standardization and validation

with tissue samples, bone DNA was amplified in 10 μl reaction mixture with 5 μl Qiagen multiplex PCR buffer, 0.2 mg/ml BSA, 0.5μM primer and 2μl of DNA. PCR conditions included an initial denaturation of 96°C for 15 minutes, followed by 40 cycles of denaturation (96°C for 30 s), annealing (50°C for 60 s) and extension (72°C for 90 s); followed by a final extension (72°C for 10 min). During all amplifications, positive extraction control and PCR negative controls were included to monitor any possible contamination. The amplicon sizes of newly designed primers ranged between 140 - 243 bps (see Supplementary Table 2 and Supplementary Figure 1 for more details on primers).

All amplified products were cleaned using Exonuclease-Shrimp Alkaline Phosphatase (GE Healthcare, USA) mixture and sequenced bi-directionally using ABI 3510xl Genetic Analyzer (Applied Biosystems Inc., USA). Sequences were aligned using MEGA v.6.0 (Tamura, Stecher, Peterson, Filipski & Kumar, 2013) and analyzed for missense or frame-shift mutations, possibly arising from sequencing errors. All sequences were then visually examined, matched against GenBank sequences and deposited in GenBank (Accession numbers: MK986797-MK986817).

2.4 Data Analyses

Data analyses were performed independently on two dataset alignments:

A) *Global dugong alignment* of 537 sequences (309 bp) consisting of Indian dugong sequences generated in this study (n=21) and sequences downloaded from GenBank (n=516) from previously published studies (Haile, 2008; Jayasankar et al., 2009; Bushell, 2013; Seddon et al., 2014; Blair et al., 2014; Plon, Thakur, Parr & Lavery, 2019). Only 88 sequences (out of 163) were used from Plon et al. (2019) due to ambiguous site calls. The sequences were arranged into global dugong distribution representing Pacific (Australia=383, Papua New Guinea=5, New Caledonia and Palau=1 each); south-east Asia (Thailand=56, Indonesia=7, Japan=2, Philippines

and Malaysia/Sabah=1 each); south Asia (India=24, Sri Lanka =4 and Mauritius=2); north-west Indian Ocean (Djibouti=8, Bahrain and Red sea=6 each, Egypt=5, United Arab Emirates=4 and Sudan=2); and south-west Indian Ocean (Tanzania=7, Madagascar=5, east Africa, Mozambique and Kenya=2 each and Comoros=1) regions.

B) *Indian dugong alignment* consisting of 21 sequences (789 bp) of mtDNA control region sequences generated in this study. These sequences were obtained from samples collected from Gulf of Kachchh (n=5), Gulf of Mannar (n=8), Palk Bay (n=4) and Andaman & Nicobar Islands (n=4).

2.4.1 Genetic lineage and phylogeography

The genetic lineage of Indian dugongs' vis-à-vis global dugong populations and genetic structure within Indian populations was determined using three different approaches:

- a) Genetic structure was estimated using Bayesian Analysis of Population Structure (BAPS) v 6.0. (Corander, Marttinen, Sirén & Tang, 2008) to identify the population clusters within both the *global dugong alignment* and *Indian dugong alignment* datasets. Models using spatial clustering of groups followed by admixture analysis were implemented in the program, with K values set between 1 to 15. Each value of K was then analyzed using 500 iterations and 100 burn-ins for each referenced individual per population.
- b) Median joining haplotype networkwas constructed using program PopArt v 1.7 (Leigh & Bryant, 2015) to assess the genetic lineage within the *global dugong alignment* as well as for the *Indian dugong alignment* datasets. Haplotype network calculations were carried for both analyses by assigning equal weights to all the variable sites.

c) Phylogenetic relationship of dugong populations was assessed only for the *global dugong* alignment. The best fit nucleotide substitution and partition schemes for the DNA dataset were selected using the Akaike Information Criterion (AIC; Akaike, 1974) implemented in program jModelTest (Posada, 2008). The best-fit substitution model was found to be HKY+I+G. Phylogenetic analysis implemented in program MrBayes v.3.2 (Ronquist et al., 2012) was conducted using Bayesian inference approach (Yang &Rannala, 1997). The chain length consisted of 21 million generations of Markov Chain Monte Carlo (MCMC) simulations, sampled every 1000 generations, with the first 3 million runs discarded as burn-ins. West Indian manatee, *Trichechus manatus* (Accession number: AY963860.1) was kept as an outgroup in the phylogenetic analysis. Finally, FigTree v.1.4 (http://tree.bio.ed.ac.uk/software/figtree/) was used to view and annotate the consensus phylogenetic tree. A posterior probability value ≥ 0.95 and above was considered for indicating strong relationships (Leache& Reeder, 2002).

2.4.2 Genetic differentiation among global dugong populations

Pairwise genetic differentiation values (F_{ST}) were computed in program Arlequin v 3.5 (Excoffier & Lischer, 2010) for *global dugong alignment* between the dugong population groups derived from multiple structure analysis. Statistical significance was considered with p-values < 0.05 between all the pairs of populations after Bonferroni correction for multiple tests.

2.4.3 Genetic diversity estimates, differentiation and demography of Indian dugongs

Genetic diversity estimates including number of haplotypes (H), total number of polymorphic sites (S), nucleotide diversity (π) and haplotype diversity (h) were calculated for the Indian dugong dataset (n=21, 789 bp sequence) using program Arlequin v 3.5 (Excoffier & Lischer,

2010). Tajima's D (Tajima, 1989), Fu's F_S (Fu, 1997) and R2 (Ramos-Onsins & Rozas, 2002) statistics and associated significance values were inferred using Arlequin v 3.5 (Excoffier & Lischer, 2010) and DnaSP v 6.12 (Rozas et al., 2017) to test for demographic signatures. Tajima's D values are in distinguishable from 0, if the populations are experiencing equilibrium state due to selective neutral variations; D= negative, during demographic expansion or mutational selection and; D=positive, during demographic contraction. Similarly, in the case of Fu's F_S statistics (Fu, 1997), negative Fu's F_S values are observed during demographic expansion or departure from null hypothesis of neutral selection and population equilibrium. Additionally, mismatch distributions for Indian dugong samples were computed using DnaSP v 6.12 (Rozas et al., 2017) to test whether the populations underwent demographic changes in the recent past. Mismatch distributions display right-skewed uni-modal peaks for populations undergoing expansion whereas ragged and multimodal peaks for populations in demographic equilibrium (Roger & Harpending, 1992). The distribution patterns were further assessed by quantifying the raggedness index (r, Harpending, 1994) and sum of squared deviations (SSD), tested for significance with 10,000 simulations using Arlequin v 3.5 (Excoffier & Lischer, 2010).

Pairwise genetic differentiation values (F_{ST}) were computed in program Arlequin v 3.5 (Excoffier & Lischer, 2010) between the populations derived from Bayesian clustering analysis. Statistical significance was considered with p-values <0.05 between all the pairs of populations after Bonferroni correction for multiple tests. A hierarchical analysis of molecular variance (AMOVA) was conducted using program Arlequin v. 3.5 (Excoffier & Lischer, 2010) to understand genetic variation among individuals, within and between populations as discussed earlier.

3. RESULTS

3.1 Phylogeography and genetic differentiation in dugongs

The Bayesian clustering, phylogeographic and phylogenetic analyses of the global dugong dataset revealed five major genetic clusters corresponding to a) three clusters grouped in the Pacific region (Australia, New Caledonia, Palau and Papua New Guinea); b) one cluster in southeast Asia (Thailand, Philippines, Japan, Malaysia/Sabah and Indonesia) and c) one cluster comprising of south Asia (Mauritius, Sri Lanka and India), northwest Indian Ocean (Red sea, UAE, Egypt, Sudan, Djibouti and Bahrain) and southwest Indian Ocean (East Africa, Tanzania, Madagascar, Mozambique, Comoros and Kenya) (Figure 1 & 2 and Supplementary Figure 2). While reporting our results, we now use the same nomenclature (Pacific, southeast Asia, south Asia, northwest Indian Ocean and southwest Indian Ocean) in this paper. In total, 76 haplotypes were identified from 537 dugong sequences (Supplementary Table 3). Highest numbers of unique haplotypes were identified from Pacific region (n = 39) followed by southeast Asia (n =20), south Asia (n=8), northwest Indian Ocean (n=6) and southwest Indian Ocean (n=2). Only two haplotypes were shared between these regions; one between Pacific, southeast Asia and south Asia and; another between Pacific, south Asia, northwest Indian Ocean and southwest Indian Ocean (Figure 2 and Supplementary Table 3).

The network and phylogenetic analyses indicate a strong phylogeographic structure and divergent mtDNA lineages among different dugong genetic clusters. The genetic differentiation (pairwise $F_{\rm ST}$ value) ranged between 0.09-0.66 across different regions (Table 1), indicating variable genetic connectivity among regions. Within each of the dugong distribution regions, relatively high haplotypic diversity ($h \pm {\rm SD}$) was found in south Asia (0.87 \pm 0.03), southeast

Asia (0.84 ± 0.03) and Pacific (0.81 ± 0.01) in comparison to the southwest Indian ocean (0.49 ± 0.1) and northwest Indian ocean (0.35 ± 0.11) regions. Overall nucleotide diversity $(\pi \pm SD)$ was considerably lower for all regions ranging from 0.01 ± 0.004 to 0.02 ± 0.01 signifying low intraregional variations (Table 2).

3.2 Genetic lineage, diversity and demography of Indian dugongs

All three different analytical approaches with the global dugong dataset (n=537, 309 bp sequence) confirm that the Indian dugongs belong to the south Asia- northwest India Ocean-southwest Indian Ocean genetic cluster with unique divergent mtDNA haplotypes. We identified a total of seven unique Indian haplotypes within the south Asia region, where two of them were shared with Sri Lanka (Figure 1 and Supplementary Table 3). One haplotype sampled from Andaman and Nicobar Islands was also observed within southeast genetic cluster (Figure 2).

We generated a longer 789 bp sequence for the Indian dugong samples collected in this study (n=21). A total of eight unique haplotypes were identified from these samples (Figure 3 and Supplementary Table 4). Palk bay, Gulf of Kachchh and Andaman & Nicobar Islands samples showed two haplotypes each, whereas Gulf of Mannar showed four haplotypes. One haplotype was shared between Gulf of Mannar, Palk bay and Gulf of Kachchh. Overall, three haplotypes were unique to Gulf of Mannar, two were unique to Andaman & Nicobar Islands and one each for Gulf of Kachchh and Palk bay, respectively (Supplementary Table 4). Mean haplotype diversity (h) and nucleotide diversity (h) was 0.85 ± 0.04 and 0.005 ± 0.001 , respectively (Table 3). Our Bayesian clustering analyses (BAPS) indicated three genetic signatures in the sampled areas. Gulf of Mannar has two clusters: one shared with Palk Bay and Andamans whereas the other one shared with Gulf of Kachchh (Figure 3). One sample from Andamans was genetically unique.

These three areas were genetically differentiated at significant level, with F_{ST} values ranging between 0.1-0.64 (Table 4). Further, AMOVA results showed higher within-population variance (63.41%) than among populations (36.59%), indicating population structure within the Indian dugongs (see Table 5).

Different demography analyses (Tajima's D, Fu's F and mismatch distribution) showed contrasting, but non-significant patterns of population demography across these areas. Gulf of Mannar samples showed a positive Tajima's D value (0.086, p=0.55), indicating population decline, whereas Palk Bay, Gulf of Kachchh and Andaman & Nicobar Islands samples showed negative values (-0.70, p=0.29; -0.83, p=0.10 and -1.09, p=0.11, respectively), indicating population expansion or selection. For Fu's F, all these regions showed positive values (Gulf of Mannar- 0.49, p=0.57; Palk bay- 1.09, p=0.62; Gulf of Kachchh- 2.2, p=0.82 and Andaman and Nicobar Island- 4.22, p=0.96, respectively) indicating population decline (Table 3). Mismatch distribution analyses reveal multi-modal peaks under both constant population size and population growth-decline models, indicating past demographic equilibrium (Figure 4). The observed mismatch values showed a statistically non-significant value of Harpending's raggedness index r = 0.56, p=0.88 (SSD=0.11,p=0.39, $R_2 = 0.12$), indicating a population equilibrium (Table 3).

4. DISCUSSION

This study provides the most exhaustive description of genetic groups of dugong at a global scale till date, and first in-depth investigation of genetic diversity and structure of Indian dugong populations. Given that our sampling represents ~10% of the available population estimates of dugongs from India (Sivakumar, 2013), the results are crucial for their conservation at regional

scale. Earlier work on dugong from Pacific (Tikel, 1997; McDonald, 1997; Seddon et al., 2014, Blair et al., 2014), southeast Asia (Palmer, 2004; Bushell, 2013, Blair et al., 2014) and western Indian ocean (Plon, Thakur, Parr & Lavery, 2019) provided an incomplete picture of genetic groupings due to limited samples from the central part of their global distribution (i.e. south Asia). Our findings fill this gap and show that the Indian samples are part of a single genetic cluster, comprising south Asia, northwest Indian ocean and southwest Indian ocean populations with low genetic differentiation. This pattern of genetic clustering is in accordance with earlier studies by Blair et al. (2013). However, these results show slightly different pattern from Plon et al., (2019), where Madagascar/Comoros formed a unique lineage within the western Indian region. It is also noteworthy to point out that such patterns could also be driven by use of poor quality sequence data from historical samples. We feel that addition of critical dugong samples from India helped in getting a clear picture of genetic groups within this region. Overall, the global data showed a very structured phylogeographic pattern with very limited sharing of haplotypes among the identified regions. While such pattern could arise from incomplete sampling effort across the dugong range, it could also indicate potential loss of gene flow among these regions due to fragmentation of contiguous dugong habitats (Marsh, O'shea & Reynolds III, 2011).

This study elucidates divergent mtDNA lineages of south Asian dugongs within the western Indian ocean populations. Indian dugong population genetically grouped within the south Asia region, although not genetically unique, consists of unique mtDNA haplotypes. Addition of novel mtDNA haplotypes from Indian dugong samples points towards high genetic diversity within south Asia. Two of the previously reported haplotypes from Sri Lanka (Plon, Thakur, Parr & Lavery, 2019) were found to be shared with southern part of dugong distribution in India at

Gulf of Mannar, whereas one haplotype sampled from Andaman & Nicobar Islands was observed within southeast Asian lineage. This indicates potential genetic connectivity between these populations in the recent past, and future work should focus on further fine-scale sampling for in-depth investigation. With local extinctions from Mauritius and Maldives and a highly imperiled dugong population in Sri Lanka (Marsh, Penrose, Eros & Hugues, 2002; Marsh & Sobtzick 2015), India holds the largest and potentially the last viable dugong populations within the south-Asia region thereby requiring immediate conservation attention.

Within India, we identified novel haplotypes from each sampling site i.e. Gulf of Mannar, Palk Bay, Gulf of Kachchh and Andaman & Nicobar Islands. Overall haplotype and nucleotide diversity for dugong populations in India were comparable to the Australia (McDonald, 1997; Blair et al., 2014; Seddon et al., 2014) and Thailand (Palmer, 2004; Bushell, 2013) dugong populations. Gulf of Mannar population showed higher haplotype diversity within the Indian regions. Presence of shared haplotypes across Gulf of Mannar, Palk Bay and Gulf of Kachchh suggest potential genetic connectivity among these populations. We found a new haplotype with longer sequences generated in this study (n=789 bp) when compared with earlier studies from Australia (McDonald, 1997; Blair et al., 2014; Seddon et al., 2014) and Thailand (Palmer, 2004; Bushell, 2013), suggesting that longer sequence data is required to assess genetic variation at regional/global scale. Finally, it is important to point out that our Indian dugong data shows contrasting patterns of population differentiation, where we found shared haplotypes among the sampled areas but high F_{ST} values. Further, the AMOVA analyses indicated signatures of within population structures. We feel that these contrasting patterns probably arise from low sample size from each areas and short sequences (less polymorphic sites) leading to the effects of genetic

drift. Further efforts through intensive sampling and more genetic data would clarify these genetic patterns in Indian dugong populations.

Similarly, our demography analyses with mtDNA show contrasting (but non-significant) signals across the sampled area, thus could not be used to deduce population growth or decline. While mitochondrial DNA has been used to assess demographic pattern (for example see Mizuno, Sasaki, Kobayashi, Haneda & Masubuchi, 2018 for Japanese harbor seals), it generally indicates evolutionary signals at longer time frame. Future work should focus on more systematic sampling effort and generate nuclear data (microsatellite, Single nucleotide polymorphisms etc.), which provide much clear signatures of recent population demography (Lah et al., 2016; Komoroske, Jensen, Stewart, Shamblin & Dutton, 2017).

Increasing human population in coastal areas and subsequent coastal developments has adversely affected nearshore ecosystems including seagrass meadows throughout their distribution range (Unsworth & Cullen, 2010), thereby impacting historically exploited seagrass-dependent dugong populations (Hines et al., 2012; Reynolds III & Marshall, 2012). These rapidly declining dugong populations (Marsh, O'Shea & Reynolds III, 2011), are further imperilled by the lack of reliable scientific data on their genetic status. This study addresses the gaps in knowledge on genetic status of Indian dugongs in comparison with the global dugong populations. Indian dugong populations retain global significance being the largest and genetically unique population in the south Asia region (Dugong CMS MoU). Based on these results, we recommend that Indian dugong populations should be managed as a "Conservation or Management Unit" to strengthen conservation strategies for ensuring their long-term survival. With concurrent implementation of Dugong Recovery Program at all the dugong distribution ranges in India (Sivakumar et al., 2018; Sivakumar et al., 2019), we hope that recovering dugongs through a scientifically robust

multidisciplinary research and participatory action would safeguard remnant dugong populations in the country.

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7. Conflict of Interest

The authors declare there are no competing interests.

8. Author Contribution

KSK, JAJ, SMconceived the study;SG, PVRPJ,KMM,SP,SD,RS, DK collected samples; YS and SM designed the experiments; YS performed the experiments and analyzed the data; YS, AP, SM wrote the manuscript; KSK, SM and AP reviewed the drafts; all coauthorsapproved the final draft.

9. Figure Legends

Figure 1: A) Global dugong distribution range and region-wise control region haplotype network generated from dugong sequences. Within each region, circles represent unique haplotypes, different colors represent country of sample origin, and size of the circle represents frequency.B) Population structure using Bayesian clustering (BAPS) analysis for global dugong dataset indicating a total of five clusters using a priori estimate of probable groups. The acronyms are as followed: i) SEIO: southeast Indian ocean; ii) NWIO: northwest Indian ocean; iii) SA: south Asia; iv) SEA: southeast Asia and v) PAC: Pacific regions.

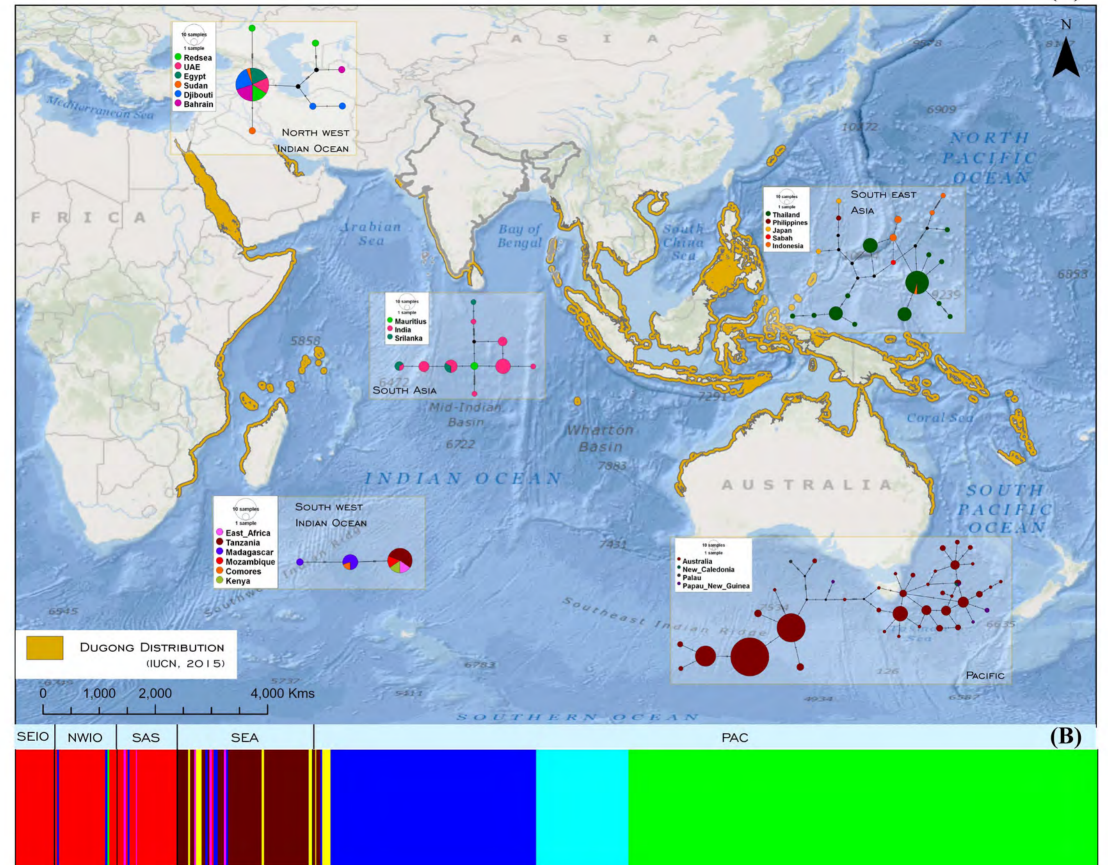
Figure 2: A) Median-joining haplotype network generated from control region sequences obtained from Indian dugong populations. Each circle represents unique haplotype, colors represent sampling locations, and size of the circle represents frequency. Donuts represent sampling locations. B) Donut colors represent proportion of unique haplotypes within each sampling location C) Population structure using Bayesian clustering (BAPS) analysis for Indian dugong dataset indicating a total of three clusters using a priori estimate of probable groups.

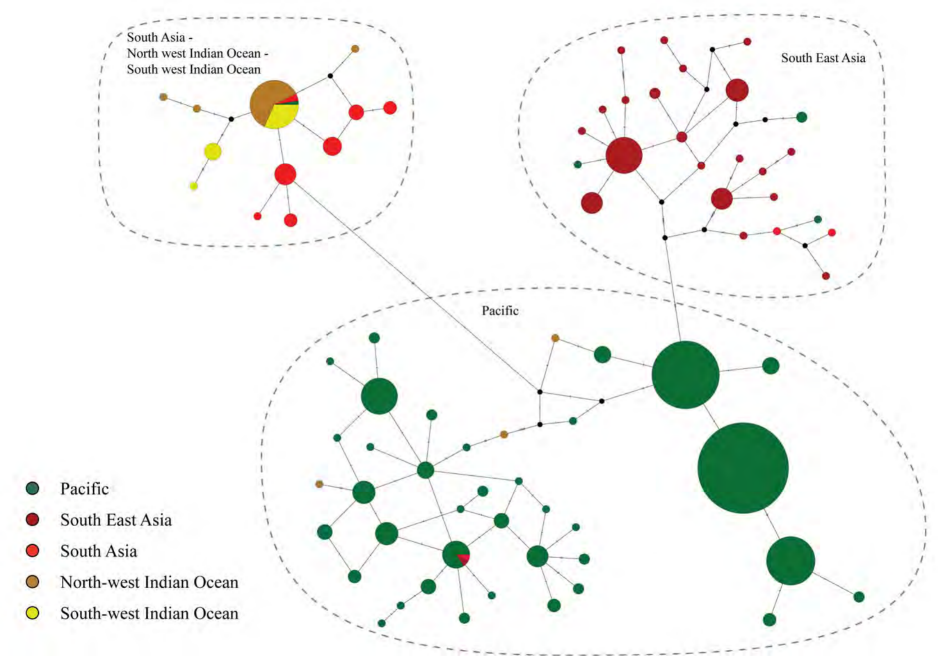
Supplementary Figure 4:Mismatch distributions of pairwise differences for Indian Dugong populations. Observed (dashed lines) and expected (solid lines) frequencies are depicted using model assuming constant population size (A) and Population growth and decline (B).

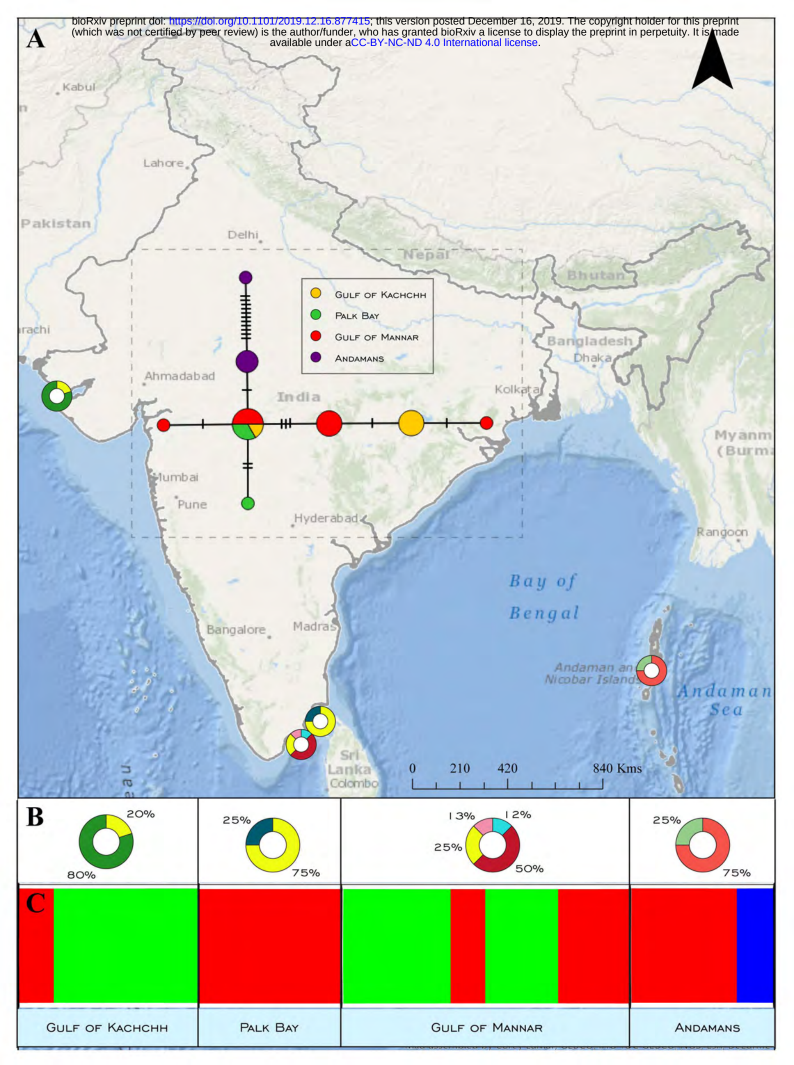
Supplementary Figure 1: Gel picture showing the lengths of amplification of designed primers with ladder on both the sides

Supplementary Figure 2: Bayesian phylogenetic tree constructed for global dugong dataset. Values at each node indicates posterior probabilities > 0.50.

Supplementary Figure 2: Median-joining haplotype network for global dugong dataset indicating three distinct population groups. Only two haplotypes are shared among these three population groups as depicted in the figure.







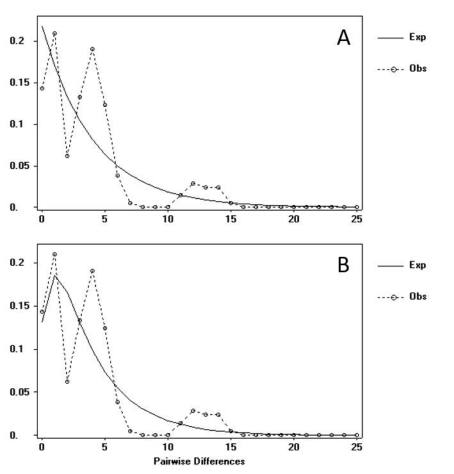


Table 1: Pairwise F_{ST} values for comparison among global dugong distribution regions

	Pacific	South East Asia	South Acia		South West Indian Ocean
Pacific	-	0	0	0	0
South East Asia	0.25*	-	0	0	0
South Asia	0.42*	0.60*	-	0	0
North West Indian Ocean	0.44*	0.66*	0.10*	-	0.01
South West Indian Ocean	0.43*	0.64*	0.16*	0.09*	-

Pairwise genetic distance values are mentioned below diagonal while above diagonal are p-values. *indicates significant differentiation values (p<0.05) after Bonferroni corrections.

Table 2: Molecular diversity estimates and demographic indices of global dugong distribution regions.

Region	Samples (n)	Haplotypes (n)	Polymorphic sites	Haplotype diversity	Nucleotide diversity	Tajima's D		Fu's Fs	
			(n)	± SD	± SD	value	p value	value	p value
Pacific	390	40	46	0.81 ± 0.01	0.02 ± 0.01	-0.106	0.547	-4.788	0.185
South East Asia	67	21	44	0.84 ± 0.03	0.02 ± 0.01	-1.077	0.129	-2.635	0.217
South Asia	30	10	25	0.87 ± 0.03	0.01 ± 0.01	-1.215	0.099	-0.249	0.487
North West Indian Ocean	31	7	20	0.35 ± 0.11	0.01 ± 0.004	-2.085	0.004	-0.408	0.448
South West Indian Ocean	19	3	7	0.49 ± 0.1	0.01 ± 0.004	0.229	0.637	3.366	0.938

Table 3: Molecular diversity estimates and demographic changes in Indian Dugong populations

Population N	N	Nh	S	h ± SD	π±SD	Tajima's D		Fu's Fs		SSD		r		R2	Reference
1 opalation	l N			11 2 0 5	11 2 00	value	<i>p</i> value	value	<i>p</i> value	value	<i>p</i> value	value	<i>p</i> value	value	Reference
Gulf of Mannar	8	4	6	0.75 ± 0.13	0.002 ± 0.002	0.086	0.55	0.49	0.57	0.01	0.95	0.08	0.90	0.17	This study
Palk Bay	4	2	6	0.50 ± 0.26	0.001 ± 0.001	-0.70	0.29	1.09	0.62	0.13	0.23	0.75	0.86	0.43	This study
Gulf of Kachchh	5	2	4	0.40 ± 0.23	0.002 ± 0.001	-1.09	0.11	2.20	0.82	0.10	0.14	0.68	0.87	0.40	This study
Andaman and Nicobar Islands	4	2	11	0.50 ± 0.26	0.006 ± 0.005	-0.83	0.10	4.22	0.96	0.20	0.26	0.75	0.90	0.43	This study
India Overall	21	8	19	0.85 ± 0.04	0.004 ± 0.00	-0.63	0.26	2.00	0.74	0.11	0.39	0.56	0.88	0.12	This study
Australia	115	52	59	0.97	2.29	-1.53 (R) -1.54 (W)		-3.81 (R) -10.01 (W)							McDonald, 2005
Australia	188	56	60	0 .94	0.02			-2.13						0.07	Blair et al.,2014
Australia	184	12		0.74 ± 0.02	0.36 ± 0.21										Seddon et al., 2014
Thailand	53	12	27	0.77 ± 0.04	1.26 ± 0.68	-0.17	0.49	1.23	0.71						Bushell 2013
Thailand	40	14		0.75 ± 0.06	0.009 ± 0.002										Palmer 2004

N=No. of samples, Nh=No. of haplotypes, S=No. of polymorphic sites, h=haplotype diversity with \pm SD, π =Nucleotide diversity with \pm SD, SSD=Sum of squared deviations

Table 4: Pairwise F_{ST} values	available under affer entration between	alling Difference populations.
51	<i>8</i>	

	Gulf of Mannar	Palk bay	Andaman & Nicobar Islands	Gulf of Kachchh
Gulf of Mannar	-	0.05	0.009	0.221
Palk bay	0.334*	-	0.029	0.041
Andaman & Nicobar Islands	0.374*	0.235*	-	0.014
Gulf of Kachchh	0.103	0.639*	0.487*	-

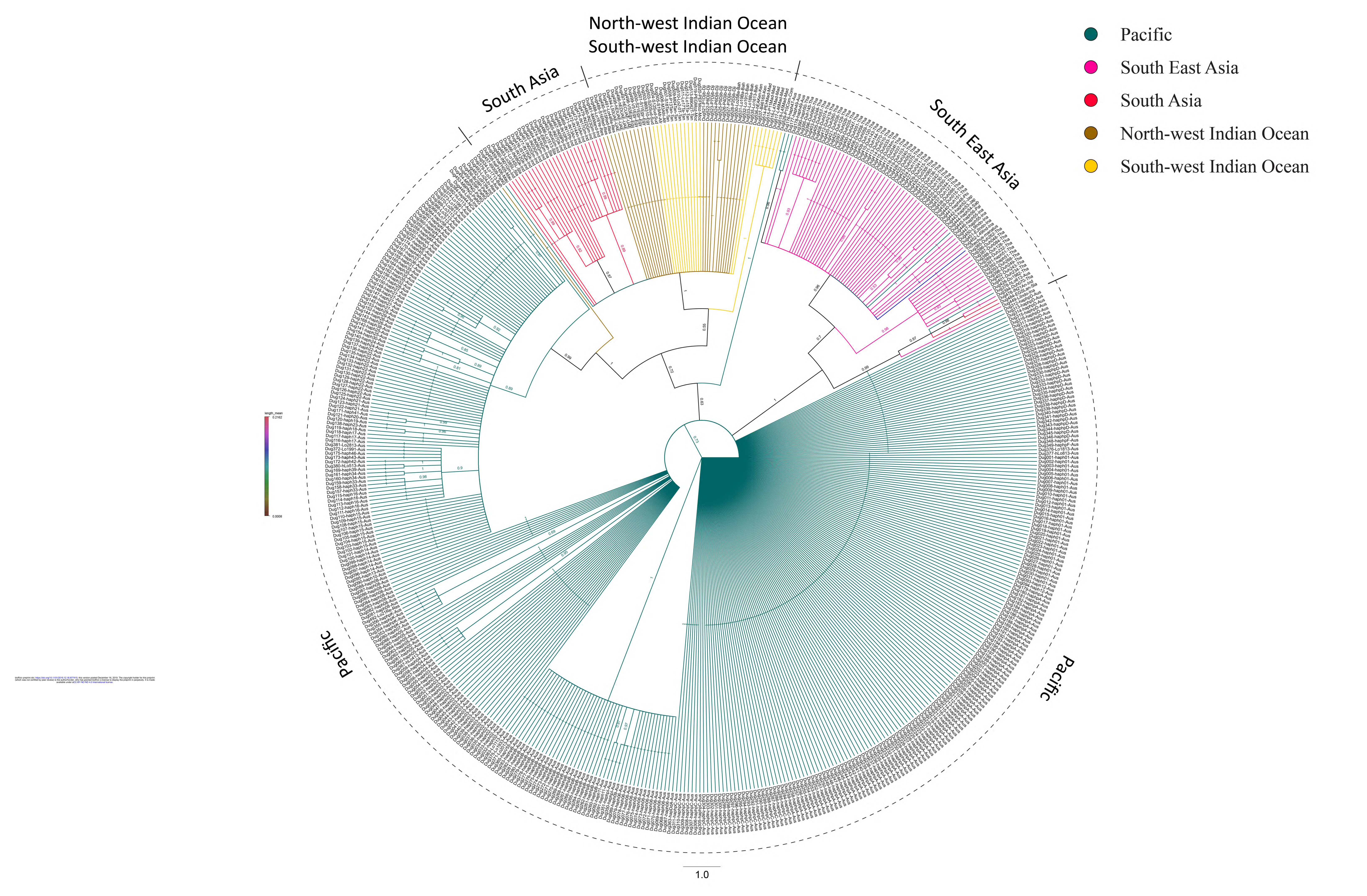
Pairwise genetic distance values are mentioned below diagonal while above diagonal are p-values.

* indicates significant differentiation values (p<0.05) after Bonferroni corrections.

 Table 5: AMOVA values for genetic structuring within and between population groups in Indian dugongs.

Structure	Source of Variation	df	Sum of Squares	Variance components	Variation (%)	Fixation index	<i>p</i> -value
All 4 nanulations grouped together	Among populations	3	14.705	0.71949 Va	36.59	φST: 0.36	0.0003
All 4 populations grouped together	Within populations	17	21.200	1.24706 Vb	63.41	ψ51.0.30	
	Among groups	2	11.121	0.35179 Va	17.27	фСТ: 0.17	0.32
Group 1: Gulf of Kachchh; Group 2: Gulf of Mannar & Palk Bay;	Among populations Within groups	1	3.583	0.43805 Vb	21.51	φSC: 0.25	0.05
Group 3: Andaman & Nicobar Islands	Within populations	17	21.200	1.24706 Vc	61.22	φST: 0.38	0.0004





Supplementary Table 1: Details of samples used in Indian dugong alignment analysis

S no.	Sample ID	Sample Type	Sampling Location	Year of Sampling	GenBank Accession Sequence ID
1	Dug-1	Tissue	Gulf of Mannar, Tamil Nadu	2017	MK986797
2	Dug-2	Tissue	Gulf of Mannar, Tamil Nadu	2017	MK986798
3	Dug-3	Tissue	Gulf of Mannar, Tamil Nadu	2017	MK986799
4	Dug-4	Tissue	Gulf of Mannar, Tamil Nadu	2017	MK986800
5	Dug-6	Tissue	Gulf of Mannar, Tamil Nadu	2017	MK986801
6	Dug-7	Bone	Gulf of Kachchh, Gujarat	2017	MK986813
7	Dug-8	Tissue	Gulf of Kachchh, Gujarat	2017	MK986814
8	Dug-9	Tissue	Palk Bay, Tamil Nadu	2017	MK986806
9	Dug-10	Tissue	Palk Bay, Tamil Nadu	2017	MK986807
10	Dug-11	Bone	Palk Bay, Tamil Nadu	2017	MK986808
11	Dug-12	Tissue	Gulf of Kachchh, Gujarat	2018	MK986815
12	Dug-13	Bone	Port Blair, Andaman & Nicobar Islands	2018	MK986809
13	Dug-14	Bone	Port Blair, Andaman & Nicobar Islands	2018	MK986810
14	Dug-15	Bone	Port Blair, Andaman & Nicobar Islands	2018	MK986811
15	Dug-16	Tissue	Gulf of Mannar, Tamil Nadu	2018	MK986802
16	Dug-17	Tissue	Gulf of Mannar, Tamil Nadu	2018	MK986803
17	Dug-19	Tissue	Gulf of Kachchh, Gujarat	2018	MK986816
18	Dug-20	Tissue	Gulf of Kachchh, Gujarat	2018	MK986817
19	Dug-21	Tissue	Shaheed Dweep, Andaman & Nicobar Islands	2018	MK986812
20	Dug-22	Tissue	Gulf of Mannar, Tamil Nadu	2018	MK986804
21	Dug-23	Tissue	Palk Bay, Tamil Nadu	2019	MK986805

Supplementary Table 2: List of primers designed to amplify the mitochondrial DNA control region in dugongs

S no.	Primer	Primer Sequence (5'-3')	Amplicon Size	Annealing Temperature
1	Dug SSF1	AAT GAA GGT CCC CGT AGT	234	50°C
1	Dug SSR1	TGC ACG ATT ATA CAT AGG	254	30 C
2	Dug SSF2	CGC GCT ATG TAC TTC GTG	203	50°C
	Dug SSR2	ATG GAC TGG ACA ATA TCC	203	30 C
3	Dug SSF3	GTA GGA TTC ATG CTC TAA	243	50°C
	Dug SSR3	GCC CGG AGC GAG AAG AGA	2-13	30 C
4	Dug SSF4	TTG ACT ACC AAG CTT CGA	217	50°C
•	Dug SSR4	CAC AGT TAT GTT ATG ATC		
5	Dug SSF5	TTC CCC TTA AAT AAG ACA	223	50°C
3	Dug SSR5	TCG AGC ATT GAC TGA ATA		30 C
6	Dug SSF6	AGG CAA ATA ACT TGT AGC	143	50°C
Ü	Dug SSR6	GCG GGA AAT GGG GT TTG		

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Haplotype ID	2	8	15	16	18	19	23	24	27	29	30	40	41	48	54	57	69	92	103	109	112	120	123	126	139	142	144	147
Hap_1	С	G	G	Т	С	Α	Т	G	C	С	Т	С	Α	Т	Т	Т	Α	Т	Α	Т	Α	С	Т	Α	С	Т	Α	С
Hap_2	•						•	•															•	•		•		
Hap_3	•			•			•		•												•		•	•		•		
Hap_4	•		•				•	•	•												•		•	•		•		
Hap_5							•		•												•		•	•		•		
Нар_6	•		•	•			•		•												•		U	•		•		
Hap_7	•		•	•			•	•	•												•		•	•		•		
Hap_8	•	•	•				C	•	•		С	Т								•	•		•	•		•		
Hap_9	•		•	•			U	•	•		С	Т									•		•	•		•		
Hap_10	•		•				U	•	•		С	Т									•		•	G		•		
Hap_11	•						•	•			С	Т											•	G		•		
Hap_12											С	Т															<u> </u>	<u> </u>
Hap_13											С	Т															<u> </u>	Ŀ
Hap_14	•	•	•				•	•	•		С	Т								•	•		•	•		•		
Hap_15											С	Т												•			<u> </u>	Т
Hap_16											С	Т												G				
Hap_17											С	Т																
Hap_18							С				С	Т												G				
Hap_19							С				С	Т												G				
Hap_20							C				С	Т												•		•		
Hap_21											С	Т												•		•		
Hap_22											С	Т												G				
Hap_23											С	Т												G				
Hap_24										Т	С	Т												G	Т	•		
Hap_25							•				С	T												G				
Hap_26							•				С	Т												•				
Hap_27	•						С				С	Т											•	G		•		
Hap_28											С	Т												G				

Haplotype																												
ID	7	3	15	16	18	19	23	24	27	29	30	40	41	48	54	57	69	92	103	109	112	120	123	126	139	142	144	147
Hap_29						Т					С	Т																
Hap_30											С	Т																
Hap_31											U	Т						C	•	•	•	•		G				
Hap_32			•	•						•	•	•	•					•	•	•	•	•						
Hap_33									•						С						G	Т	•		•	•		
Hap_34			•						•		С	Т					•	•	•	•	•	•	•	G				
Hap_35		Α						Α		•	•	•							G		G							
Hap_36											•	•							•	•	•	•						
Hap_37			•	•						•	U	•	•		С			•	•	•	G	•						
Hap_38											U	•		•	С				•	•	G	•						
Hap_39											•	•							•	•	G	Т				С		
Hap_40			•							•	•	•	•	•				•	G	•	G	•						
Hap_41											•	•		•					•	•	G	Т				С		
Hap_42										•	•	•									G	Т				С		
Hap_43			•							•	U	•	•	•	С			•	•	•	G	•						
Hap_44										•	U	•	•		С					•	G	•		•				
Hap_45												Т									G	Т				С		
Hap_46																					G	Т				С		
Hap_47			Т	G	G					•	•	Т	G			G	G			G	G	Т		•		C		
Hap_48			•	•						•	•	•	•					•	•	•	G	Т				C		
Hap_49		•								•	U	•		•	С				•	•	G	•		•		С		
Hap_50											•	•							•	•	G	Т				С		
Hap_51			•	•						•	•	•	•	Α				•	•	•	G	Т		•		C		
Hap_52									Т		•	•									G	Т						
Hap_53																					G	Т				С		
Hap_54										•		•									G	Т				С		
Hap_55																					G	Т				С	G	
Hap_56																			G	•	G	•						
Hap_57		•	•					•	•		•	•	•	•			•	•		•	G	Т	•	•		С		

Haplotype																		, .	3	6	7	0;	3	97	69	7	4	7.
ID	7	3	15	16	18	19	23	24	27	29	30	40	41	48	54	57	69	92	103	109	112	120	123	126	139	142	144	147
Hap_58											С	Т												G				
Hap_59											С	Т										•		G				
Hap_60																												
Hap_61							С																					
Hap_62																												
Hap_63																												
Нар_64																												
Hap_65																			G									
Нар_66																												
Нар_67																			G		G							
Нар_68																			G		G							
Hap_69																												
Hap_70							С				С	Т																
Hap_71																						Т						
Hap_72																												
Hap_73	Т																											
Hap_74																								G				
Hap_75																								G				
Нар_76												•										•		•		•		

List of mitochondrial control region haplotypes and variable sites in global dugong dataset

Hanlatyna										Ī		<u> </u>								l								
Haplotype ID	151	152	153	154	156	157	160	161	163	166	169	171	172	173	174	177	178	179	185	188	195	199	200	207	208	209	216	225
Hap_1	Α	Α	Т	Α	С	С	С	Т	С	С	Т	С	Α	Α	С	С	С	Α	Т	Α	С	Т	С	С	С	Т	С	Т
Hap_2																												
Hap_3	G																											
Hap_4																			•				•		Т	С		
Hap_5																							•					С
Нар_6							•												•				•		Т	C		
Hap_7										Т									•				•		Т	С		
Hap_8		G			Т	Т	Т	С		Т											•	•	•					
Hap_9		G			Т	Т	Т	С		Т							Т		•				•					
Hap_10		G			Т	Т	Т	С		Т							Т		•				•					
Hap_11		G			Т	Т	Т	С		Т							Т						•		Т			
Hap_12		G			Т	Т	Т	С		Т							Т		•				•					
Hap_13		G			Т	Т	Т			Т									•				•					
Hap_14		G			Т	Т	Т	С		Т						Т	Т						•					
Hap_15		G			Т	Т	Т	С		Т							Т		•				•					
Hap_16		G			т	Т	Т	С		Т							Т									•	•	
Hap_17		G			Т	Т	Т	С		Т																		
Hap_18		G			Т	Т	Т	С		Т							Т							Т				
Hap_19		G			Т	Т	Т	С		Т	С						Т											
Hap_20		G			Т	Т	Т	С		Т							Т							Т				
Hap_21		G			Т	Т		С		Т							Т											
Hap_22		G			Т	Т		С		Т							Т		•				•					
Hap_23		G			Т	Т	Т	С		Т							Т								Т	С		
Hap_24		G			Т	Т	Т	С		Т							Т								Т			
Hap_25		G			Т	T		С		Т			•				T											
Hap_26		G			Т	Т	Т	С		Т							Т											
Нар_27		G			T	T	T	С		T							T										•	
Hap_28		G			Т	T	Т	С		Т											Т		•		Т			

Haplotype										1.0																	, ,	
ID	151	152	153	154	156	157	160	161	163	166	169	171	172	173	174	177	178	179	185	188	195	199	200	207	208	209	216	225
Hap_29		G			Т	Т	Т	С		Т																		
Hap_30		G			Т	Т	Т	С		Т							Т								Т			
Hap_31		G			Т	Т	Т	С		Т							Т								Т			
Hap_32							Т			Т																		
Hap_33				G	Т			С										G										
Нар_34		G			Т	Т	Т	С		Т							Т		-									
Hap_35		G				Т		С	Т				G															
Hap_36		G			Т	Т	Т	С		Т							Т											
Hap_37						Т		С	Т			Т															Т	
Hap_38						Т		С	Т			Т											Т				Т	
Hap_39			С	G	Т			С	Т									G										
Hap_40		G						С	Т				G		Т													
Hap_41			С	G	Т			С	Т																			
Hap_42			С	G	Т			С	Т														Т					
Hap_43	•					Т	•	С	Т	•	•	Т	•	•	•		•	•		•	•	•			Т		Т	
Hap_44	•					Т		С	Т			Т													Т		G	
Hap_45	•		С	G	Т	•		С	Т		•	•	•		•		•	•										
Hap_46	•		С	G	Т	•	•	С	Т	•	•	•	•	•	•		•	•		Т	•	•						
Hap_47			С	G	Т	•	•	С	Т		•	•	•		•					•	•	•						
Hap_48		•	C	G	Т	•	•	C	Т	•	•		•		•		•	•	•	•	•	•				•		
Hap_49	•					Т	•	С	Т		•	Т	•	•	•		•	G		•	•	•			•		Т	
Hap_50			С	G		•	Т	С	Т		•	•	•		•					•	•	С						
Hap_51		•	C	G		•	Т	U	Т	•	•		•		•				•	•	•	C			•	•		
Hap_52	•		С	G		•	Т	С	Т	•	•	•	•	•	•		•	G		•	•	•			•			
Hap_53	•			G	Т			С	Т																			
Hap_54			С	G	Т			С	Т		•	•	G											•				
Hap_55			С	G	Т			С	Т																			
Hap_56		G				T		С	Т																			
Hap_57			С	G	Т			С	Т																			

Haplotype	3,1	152	153	154	156	157	160	51	163	166	169	1	72	173	174	177	178	179	185	188	195	199	200	7	808	209	216	35
ID	151		15	15				161	16		16	171	172	17	17	17		17	18	18	15	15	20	207	20	20	21	225
Hap_58		G			Т	Т	Т	С		Т			G				Т											
Hap_59		G			Т	Т	Т	С		Т				G			Т											
Hap_60			С			Т				Т			•	G	Т		•							•				
Hap_61			С			Т				Т				G	Т				С									
Hap_62			С			Т				Т				G	Т				С									
Hap_63			С							Т				G	Т													
Hap_64			С			Т				Т				G	Т				С									
Hap_65			С							Т				G	Т													
Hap_66			С							Т				G	Т								Т					
Hap_67		G				Т		С	Т				G		Т													
Hap_68		G				Т		С	Т				G															
Hap_69														G														С
Hap_70		G			Т	Т	Т	С		Т				G			Т											
Hap_71			С			Т				Т				G	Т													
Hap_72	G					Т				Т				G														
Hap_73	G				•	Т				Т				G														
Hap_74		G	С			Т				Т				G	Т													
Hap_75		G	С			Т				Т				G	Т													
Hap_76		G			Т	Т	Т	С		Т				G					•	•	•							

			r										r			
Haplotype ID	230	231	234	235	240	253	256	268	271	273	622	291	967	297	Source	Region
Hap_1	Т	С	Т	С	Α	С	С	С	Т	Т	С	С	G	G	Australia (143)	Pacific
Hap_2			С												Australia (79)	Pacific
Hap_3			С												Australia (5)	Pacific
Hap_4															Australia (41)	Pacific
Hap_5		•	С				•	•	•	•	•	•	•		Australia (5)	Pacific
Hap_6															Australia (3)	Pacific
Hap_7		•	•						•		•	•	•	•	Australia (2)	Pacific
Hap_8		•	С		G		•	Т	С	•	•	•	•	Α	Australia (1)	Pacific
Hap_9			С		G			Т	С					Α	Australia (9)	Pacific
Hap_10		•	C		G			Т	C	•	•	٠	•	Α	Australia (9)	Pacific
Hap_11		•	C					Т	C	•	•	•	•	Α	Australia (8)	Pacific
Hap_12		•	C		G			Т	C	•	•	•	•	Α	Australia (5)	Pacific
Hap_13		•	U		G			Т	U	•	•	•	•	Α	Australia (2)	Pacific
Hap_14		•	C		G			Т	C		•			Α	Australia (1)	Pacific
Hap_15			С		G			Т	С					Α	Australia (2)	Pacific
Нар_16			С		G	-		Т	С	•	•	•		Α	Australia (11), Indonesia (1), India (1)	Shared haplotype
Hap_17		•	U		G			Т	U		•		•	Α	Australia (23)	Pacific
Hap_18		•	С		G			Т	С					Α	Australia (3)	Pacific
Hap_19		•	U					Т	U		•		•	Α	Australia (2)	Pacific
Hap_20			С		G			Т	С					Α	Australia (4)	Pacific
Hap_21	С	•	С		G			Т	С					Α	Australia (1)	Pacific
Hap_22	С		С		G			Т	С					Α	Australia (1)	Pacific
Hap_23		•	C					Т	С					Α	Australia (3)	Pacific
Hap_24			С					Т	С					Α	Australia (2)	Pacific
Hap_25			С		G			Т	С					Α	Australia (4)	Pacific
Hap_26		•	С					Т	С					Α	Australia (1)	Pacific
Hap_27			C					Т	C					Α	Australia (1)	Pacific
Hap_28			С				•	Т	С				•	Α	Australia (2)	Pacific

Haplotype ID	230	231	234	235	240	253	256	897	271	273	279	291	296	297	Source	Region
Hap_29			С		G			Т	С					Α	Australia (1)	Pacific
Hap_30			С					Т	С					Α	Australia (1)	Pacific
Hap_31			С					Т	С					Α	Australia (1)	Pacific
Hap_32			С		G										Australia (1)	Pacific
Hap_33		Т	С		G			Т	С						Australia (2)	Pacific
Hap_34	•		С					Т	С					Α	Australia (2), New Caledonia (1), Papau New Guinea (1)	Pacific
Hap_35			С						С						Australia (1)	Pacific
Hap_36			C		G			Т	С	•	•	•		Α	Australia (1)	Pacific
Hap_37		Т	U		•	•	•	T	С	•	•	•	Α	•	Thailand (8)	South East Asia
Hap_38		Т	U		•	•	•	Т	С	•	•	•	Α		Thailand (1)	South East Asia
Hap_39		Т	C		G			Т	С	•	•	•			Thailand (9)	South East Asia
Hap_40			U		•	•	•	•	С	•	•	•	•	•	Thailand (1)	South East Asia
Hap_41			U		G	•	•	Т	С	•	•	•	•		Thailand (22), Indonesia (1)	South East Asia
Hap_42			C		G			Т	С	•	•	•			Thailand (8)	South East Asia
Hap_43	•	Т	C		•	•	•	Т	С	•		•	Α		Thailand (1)	South East Asia
Hap_44	•	Т	С		•	•	•	T	С	Α	•	G	Α	•	Thailand (1)	South East Asia
Hap_45	•		C		G	•	•	Т	С	•		•	•		Thailand (1)	South East Asia
Hap_46	•	•	U		G	•	•	Т	C	•		•	•	•	Thailand (1)	South East Asia
Hap_47	•		C		G			Т	С			•	•		Thailand (1)	South East Asia
Hap_48	•	•	U		G		G	Т	C	•		•	•	•	Thailand (1)	South East Asia
Hap_49		Т	U		G	•	•	Т	С	•	•	•	Α	•	Thailand (1)	South East Asia
Hap_50		Т	С		G			Т	С						Philippines (1)	South East Asia
Hap_51		Т	U		G	•	•	Т	С	•	•	•	•	•	Japan (1)	South East Asia
Hap_52		Т	U		G	•	•	Т	С	•	•	•	•	•	Japan (1)	South East Asia
Hap_53		Т	С		G			Т	С						Sabah (1)	South East Asia
Hap_54			С		G			Т	С						Palau (1)	Pacific
Hap_55		Т	С		G			Т	С						Indonesia (2)	South East Asia
Нар_56			С						С						Indonesia (1)	South East Asia
Hap_57		Т	С		G			Т	С						Indonesia (2)	South East Asia

Haplotype ID	230	231	234	235	240	253	256	897	271	273	279	291	967	297	Source	Region
Hap_58			C		G			T	С		•	•	•	Α	Papau New Guinea (1)	Pacific
Hap_59			С		G			Т	С					Α	Papau New Guinea (2)	Pacific
Hap_60			С		G			Т							Papau New Guinea (1), Mauritius (2), Red Sea (4), UAE (4), Egypt (5), Sudan (1), Africa (2), Tanzania (7), Mozambique (2), Djibouti (6), Bahrain (5),	Shared haplotype
Hap_61		Т	С		G			Т							Kenya (2) India (1), Srilanka (2)	South Asia
Hap_62	٠	'	С	•	G	•	•	T	٠	٠	•	٠	•	٠	India (4), Sri lanka (2)	South Asia
Hap_63			С		G			T							India (8)	South Asia
Hap_64		Т	С		G			Т							India (4)	South Asia
Hap_65			С		G			Т							India (3)	South Asia
Hap_66			С		G			Т							India (1)	South Asia
Hap_67			C									•	•		India (1)	South Asia
Hap_68			С	•	•			•	С		•	•	•		Sri lanka (1)	South Asia
Hap_69			C		G			•			•	•	•		Red Sea (1)	North West Indian Ocean
Hap_70			U		G			Т		•		•	•	•	Red Sea (1)	North West Indian Ocean
Hap_71		Т	С		G			Т							Sudan (1)	North West Indian Ocean
Hap_72			С	Т	G			Т							Madagascar (4), Comores (1)	South West Indian Ocean
Hap_73			С	Т	G	Т		Т			Т				Madagascar (1)	South West Indian Ocean
Hap_74			C	Т	G			Т						Α	Djibouti (1)	North West Indian Ocean
Hap_75			С	Т	G			Т							Djibouti (1)	North West Indian Ocean
Hap_76			C		G			Т	С			•	•		Bahrain (1)	North West Indian Ocean

 Table 2: Control region haplotypes and variable sites in Indian dugongs

Haplotype ID									Site	num	bers									Source	Global ID
	33	32	136	216	225	265	266	270	274	276	279	285	286	298	313	344	353	381	456		
Hap_1	Т	Т	С	Α	Α	Α	С	Т	Т	С	Т	Α	G	С	С	Т	G	Т	G	Gulf of Mannar (1)	Hap_61
Hap_2			Т													С				Gulf of Mannar (4)	Hap_62
Hap_3			Т					С						Т		С			Α	Gulf of Mannar (2), Palk Bay (3), Gulf of Kachchh (1)	Hap_63
Hap_4			Т					С						Т	Т	С			Α	Gulf of Mannar (1)	Hap_66
Hap_5	С	С	Т					С						Т		С			Α	Palk Bay (1)	Hap_77
Нар_6			Т	G				С						Т		С			Α	Andaman & Nicobar Islands (3)	Hap_65
Hap_7	-		Т	G	G	G	Т	-	С	Т	С	G	Α	Т		С	Α	С	Α	Andaman & Nicobar Islands (1)	Hap_67
Hap_8	-		Т									-								Gulf of Kachchh (4)	Hap_64

Numbers mentioned above each nucleotide base represents positions relative to the common 789 bps sequence used for comparison.