Molecular Assessments, Statistical Effectiveness Parameters and Genetic Structure of Captive Populations of *Tursiops truncatus* Using 15 STRs

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**Abstract:** Genetic analysis is a conventional way of identifying and monitoring captive and wildlife species. Knowledge of statistical parameters reinforcing their usefulness and effectiveness as powerful tools for preserving diversity is crucial. Although several studies have reported the diversity of cetaceans such as *Tursiops truncatus* using microsatellites, its informative degree has been poorly reported. Furthermore, the genetic structure of this cetacean has not been fully studied. In the present study, we selected 15 microsatellites with which 210 dolphins were genetically characterized using capillary electrophoresis. The genetic assertiveness of this set of hypervariable markers identified one individual in the range of $6.927 \times 10^3$ to $1.806 \times 10^6$, demonstrating its substantial capability in kinship relationships. The genetic structure of these 210 dolphins was also determined regarding the putative capture origin; a genetic stratification ($k = 2$) was found. An additional dolphin group of undetermined origin was also characterized to challenge the proficiency of our chosen markers. The set of markers proposed herein could be a helpful tool to guarantee the maintenance of the genetic diversity rates in conservation programs both in *Tursiops truncatus* and across other odontocetes, Mysticeti and several genera of endangered and vulnerable species.

**Keywords:** *T. truncatus*; conservation programs; microsatellites; gene-flow; dolphins; population genetics
1. Introduction

Delphinidae (dolphins and killer whales) is a diverse family including at least 12 genera and 37 species [1]. The bottlenose dolphin (Tursiops truncatus, Montagu, 1821) is among the most mobile species, presenting a broad distribution in the different oceans and polar coastal waters [2]. This species exhibits size variation related to feeding, with a length ranging from 2.5 m to 3.8 m [3]. Regarding the depth of its dives, it depends on the hour, being greater than 450 m at night and tending to be shallow (50 m) during the day [4].

*T. truncatus* has been characterized depending on its distribution in coastal (>20 m isobath), continental shelf (between 20–200 m depth), inshore (i.e., bays, estuaries, and sound), and oceanic (>200 m depth) waters [5]. Each assignment shows morphological, demographic, spatial, temporal, and genetic differences [6]. Given its social capacity, *T. truncatus* is one of the most used animals in the recreational industry and assisted therapy, being one of the most widely distributed specimens in captivity [7]. In addition, research programs have included specimens living as captives to enrich the knowledge about their social lives, communication, cognitive development, natural ecology, and feeding habits, among others [8–10].

In agreement with the Mexican “Ley General de Vida Silvestre” (article 60 Bis), any species of marine mammal must not be subject to extractive use, whether for subsistence or commercial purposes [11]. Nonetheless, the Mexican aquariums and zoos still house many dolphins. Of note, this law did not prohibit scientific research and education programs, providing a unique opportunity to understand their physiology, cognitive capacities, and risk of illness [9,12–14]. The research and educational programs have generated scientific and robust information about the reproductive biology of these and other cetaceans with significant benefits for the conservation of wild populations [9]. Such investigations have facilitated reproductive assistance through biobank generation, which could help prevent the extinction of vulnerable species [9]. However, genetic and phenotype diversity maintenance is imperative to ensure the adaptive potential and persistence (long-term) in conservation and sustainability [15].

Despite its low-cost, morphological identification could be a subjective feature. To promote the most effective protection, the characterization of the genetic population architecture (diversity and structure) is crucial for efficient and effective preservation practices [6]. Thus, the first step in the reintroduction programs is to determine the genetic structure, especially in those species lacking physical dispersal barriers and coming from different geographic origins [16,17]. Thus, genetic tools are cornerstones that highlight the reproductive potential and delineate the diversity and phylogenetic relationships [18].

Microsatellites or short tandem repeats (STRs) are multiallelic markers appropriate for population genetic studies and pedigree analyses. These hypervariable molecular markers may be amplified with modest DNA concentrations, being a cheaper process than next-generation sequencing methods [19]. However, the informativeness degree and the set number of decision-making markers supporting the conservation programs and protecting biodiversity have been poorly studied. Mainly, the parentage identity allows for identifying genetic differentiation levels, a fundamental aspect of reproduction strategies' success [20,21]. Although many articles support the use of STRs, only a limited number have reported the probability of identity [22,23].

In this study, the genetic diversity and population structure of 210 captive dolphins (*T. truncatus*) from 18 Mexican dolphinariums were characterized using 15 STRs. This set of markers allowed us to genetically distinguish each dolphin with remarkable efficiency (discrimination power $\geq 0.999$), evidencing its robustness at a low cost. Our findings validated using these 15 STRs to evaluate the probability of identity and characterize the genetic structure and diversity of genome banks used in artificial insemination, which has been poorly described.
2. Materials and Methods

2.1. Study Population

The present study was conducted in agreement with the Federal Attorney for Environmental Protection (PROFEPA, initials in Spanish) in Mexico City to carry out joint actions for the diagnosis, surveillance, prevention, and control of diseases in any animal species whose survival is threatened. The study was approved by the Research Committee of the Escuela Superior de Medicina from the Instituto Politécnico Nacional (ESM-IPN, initials in Spanish).

Field research was carried out in 18 Mexican centers of dolphin captivity, distributed in eight states with tourist impact (Supplementary Materials Table S1 and Figure S1). Two hundred and ten blood samples were obtained from the caudal fins of *T. truncatus* living captive. The capture locality was collected in 177 individuals from expedients, grouping these specimens according to this information. Nevertheless, its demographic history was unknown. Out of 210 dolphins, 104 were putatively assigned to Mexican waters (MD; 92 of these were obtained from the California Gulf, MD1, and 12 from the Gulf of Mexico, MD2). Likewise, 46 dolphins were assumed to be from Cuban territorial waters (CD), 14 from southern Australian waters (AD), and 5 from Japanese (JD) waters. The rest of the dolphins (41) had an unknown geographic origin (undetermined dolphins; UD) (Supporting information Table S1).

2.2. Genotyping Analysis

Genomic DNA was isolated using the MagNA Pure LC Total Nucleic Acid Isolation kit (Qiagen, Hilden, NRW, Germany) in the MagNA Pure LC automated equipment (Roche, Germany). DNA purity ($\lambda_{260}/\lambda_{280}$) and concentration (ng/µL) were evaluated with NanoDrop 1000 (Thermo Fisher Scientific, Suwanee, GA, USA). DNA integrity was checked with agarose (Sigma Aldrich, St. Louis, MO, USA) gel 0.8% stained with ethidium bromide (Sigma Aldrich, St. Louis, MO, USA).

Samples were genotyped with 15 microsatellite loci previously reported (Table 1). These 15 STRs were chosen based on available information from earlier population genetic studies, and an observed heterozygosity of at least 0.600 was considered [24–29]. PCR was performed in a total volume of 10 µL containing 200 nM of each primer, 35 ng DNA template, 200 mM of dNTPs, 1 × reaction buffer, 1.5 mM of MgCl₂, and 0.03 U of Taq DNA polymerase (Roche Diagnostics GmbH, Mannheim, BW, Germany). The amplification conditions consisted of 35 cycles of an initial denaturation at 94 °C for 3 min, followed by a denaturation at 94 °C, annealing (depending on each primer; Table 1) and extension at 72 °C for 1 min. Thermal cycling was carried out on an Applied Biosystems Thermal Cycler (Veriti 96 Well Thermal Cycler, Carlsbad, CA, USA). Primers at the 5’- end of forward were labelled with the 6-FAM fluorescent dye. Allelic discrimination was carried out in a 3730xl DNA analyzer (Applied Biosystems, Carlsbad, CA, USA). Capillary electrophoresis conditions were 15 kV at a constant temperature of 60 °C with POP-7 polymer for 24 min (Applied Biosystems, Carlsbad, CA, USA); GeneScan™ 500 TAMRA™ dye was used as the internal size standard (Applied Biosystems, Carlsbad, CA, USA). Allelic assignment was performed with the GeneScan v3.2 software (Applied Biosystems, Carlsbad, CA, USA).
Table 1. Sequence of 15 loci explored in the four putative groups of *T. truncatus*.

| Locus   | Gene Bank Access Number | Tandem Repeat | Sequence primer (5′–3′)       | Allele Size | Annealing T | Reference |
|---------|-------------------------|---------------|--------------------------------|-------------|-------------|-----------|
| D08     | NA                      | (TG)n         | GATCCATCATATTGTCAAGTT          | 94–122      | 58          | [27]      |
|         |                         |               | TCCCTGGGTTGAIAGTCTTC          |             |             |           |
| EV37    | NA                      | (AC)n         | AGCTTGAATTGTGAAGTCAAGTT     | 189–241     | 56          | [28]      |
|         |                         |               | TAGTAGAGCCGTGAAAAGGCTC       |             |             |           |
| KWM2    | NA                      | (AC)n         | GCTGTGAAAATATTTAATGTT      | 138–160     | 47          | [29]      |
|         |                         |               | CACTGTGGCACAAATGTTGA         |             |             |           |
| KWM9    | NA                      | (AC)n         | TGTCACCCAGGAGGACAAGG        | 170–188     | 59          | [29]      |
|         |                         |               | GGGAGGGGAGGAGTTTCTG         |             |             |           |
| KWM12   | NA                      | (AC)n         | CCATCAATCCACGCTGCTC         | 186–160     | 50          | [29]      |
|         |                         |               | CACTGCAGAATGATGACC          |             |             |           |
| MK6     | AF237891                | (GT)n         | GCTCTCTTTCCAGGTGTAAGCC      | 147–187     | 51          | [24]      |
|         |                         |               | GCCCCACTAATTGTGTTGCAACC     |             |             |           |
| MK8     | AF237892                | (CA)n         | TCCTGGAGCATCTTATAGTGCC      | 80–114      | 58          | [24]      |
|         |                         |               | CTCTTGACATGCCCTTACC         |             |             |           |
| MK9     | AF237893                | (CA)n         | CAAACCAAGTGAGGACTCCC        | 161–175     | 54          | [24]      |
|         |                         |               | TTATCCCTTGTTGCTGACGTG       |             |             |           |
| Ttr04   | DQ018982                | (CA)n         | CGGACCCGACTTTCCACG          | 103–127     | 65          | [24]      |
|         |                         |               | GTTGTGTCCCCAGGATTTGAC       |             |             |           |
| Ttr11   | DQ018981                | (CA)n         | CTTGCAACTGCCTTCTGGG         | 193–219     | 61          | [25]      |
|         |                         |               | GCTTTGCCACATCAAAGGGAGTGA    |             |             |           |
| Ttr19   | DQ018980                | (CA)n         | TGGGTGGACCTTACAAAATCG       | 182–200     | 54          | [24]      |
|         |                         |               | GTTTAAGGGCTGTAAGAGG         |             |             |           |
| Ttr58   | DQ018985                | (CA)n         | TGGGTCTTGAGGGGTCTGG        | 166–194     | 62          | [25]      |
|         |                         |               | GTTGGCTGGGAGGCTCTTGTTG      |             |             |           |
| Ttr63   | DQ018986                | (CA)n         | CAGCTTACAGGCAAATGAGAG       | 83–149      | 59          | [25]      |
|         |                         |               | GTTTCCCTATGCGAGGATCATCA     |             |             |           |
| TexVet5 | AF004905                | (CA)n         | TATGTGCTAAATGGAGACAG        | 196–216     | 55          | [26]      |
|         |                         |               | TTGAGATGACTCTGCTGCTGGG      |             |             |           |
| TexVet7 | AF004907                | (CA)n         | TGCACCTGAGGGTGTCAGGAG      | 155–169     | 64          | [26]      |
|         |                         |               | CTTAATTGGGGCGATTTCAC        |             |             |           |

Note: NA = Not available in the Gene Bank database.

2.3. Statistical Analysis

The obtained genotypes were subjected to statistical analyses, including allele and genotype frequencies, number of alleles (k), and expected (He) and observed (Ho) heterozygosity using Arlequin v3.5.2.2; (Berne, Switzerland) [30]. The number of effective alleles (Ne) and Shannon’s index (I) were determined with GenAlEx v6.5; (Canberra, Australia) [31]. Hardy–Weinberg’s (HW) expectation by Weir and Cockerham’s F statistic (FIS) was estimated with Genetix v4.05.2; Genetix team (Montpellier, France) using 10,000 permutations. Linkage disequilibrium (LD) was estimated with FSTAT v2.9.3.2; (Lausanne, Switzerland) using 105,000 permutations [32].

Null alleles and large dropouts were checked for each locus with Genepop v4.6 and FreeNA and confirmed in Micro-Checker v2.2.3; (Hull, UK) [33,34].

Matching probability (MP), discrimination power (PD), and the polymorphic information content (PIC) were assessed using PowerStats v1.2; Promega CORP.(Wisconsin, USA) [35].

The genetic structure was inferred with Structure software v2.3.3; (Oxford, UK) using 100,000 burn-in and 700,000 after-burn-in interactions [36]. Five independent runs were
performed for each K-value (1 to 10). The output file was analyzed to find the most probable K-value with the Structure Harvester program [37]. The genetic distances ($F_{ST}$ values) were assessed with Arlequin v3.5.2.2; (Berne, Switzerland) (10,000 permutations) and adjusted by a false discovery rate test in R-software; (Tel Aviv, Israel) [30,38]. These values were visualized in a multidimensional scale plot (MDS) with SPSS v11; IBM CORP. (New York, NY, USA) [39].

3. Results

3.1. Microsatellite Diversity Parameters and Paternity Effectiveness

Overall, all loci presented at least seven different alleles with a mean PIC of 0.766. The most informative markers were Ttr63 ($k = 26$), EV37 ($k = 24$), and MK06 ($k = 18$), whereas TEXVET07 ($k = 7$), MK09 ($k = 8$), Ttr58 ($k = 8$) and KWM02 ($k = 9$), were the least informative (Supplementary Materials Table S2). Of note, the allele frequency distributions exhibited bi- and trimodal distributions.

MP and PD are summarized in the Supplementary Materials Table S2. Even though some loci individually showed low diversity, globally, all 15 STRs were highly informative, identifying one individual in billions of dolphins ($1.141 \times 10^{-16}$) with a highly efficient PD (PD $\geq 0.999$).

3.2. Null Alleles, Hardy–Weinberg Expectation, and Linkage Disequilibrium

Given the high inbreeding values ($F_{IS}$ mean = 0.198; range: 0.058–0.423), we determined and confirmed null alleles using two different programs. The null allele frequencies ranged from 0.023 to 0.185 with wide confidence intervals (Supplementary Materials Table S3), except in Ttr11, Ttr58, and TEXVET5. None of these loci presented evidence of large allele dropout.

Regarding the HW expectations, we found a remarkable departure (HWD) in 13 out of 15 loci even after Bonferroni’s correction ($p \leq 0.0034$). The HWD was related to a homozygosis excess ($F_{IS} > 0$). A significant number of loci pairs exhibited LD adjusting the $p$-value for 5% ($p \leq 0.00047$) and 1% ($p \leq 0.000095$) of the nominal level (Supplementary Materials Table S4).

3.3. Population Structure

$F_{ST}$ values were also estimated to compare the variation among the populations; UD was included in this analysis (Table 2). High population differentiation was presented between all groups, showing them well-separated from each other. The most separated dolphin groups were CD vs. JD, followed by CD vs. AD. Lowest but significant differences were found between CD and MD; MD and UD showed a nuance genetic distance and in turn, a nonsignificant genetic difference.

Table 2. $F_{ST}$ values estimated from 15 loci studied in the four groups of T. truncatus.

|        | AD  | CD  | JD  | MD  | UD  |
|--------|-----|-----|-----|-----|-----|
| AD     | -   | $\leq 0.0001$ | $\leq 0.0001$ | $\leq 0.0001$ | $\leq 0.0001$ |
| CD     | 0.27886 | -   | $\leq 0.0001$ | $\leq 0.0001$ | $\leq 0.0001$ |
| JD     | 0.15746 | 0.23717 | -   | $\leq 0.0001$ | $\leq 0.0001$ |
| MD     | 0.17679 | 0.05293 | 0.11988 | -   | $0.25225 \pm 0.0264$ |
| UD     | 0.20180 | 0.03210 | 0.13256 | 0.00965 | -   |

Note: AD: Australian dolphins; CD: Cuban dolphins; JD: Japanese dolphins; MD: Mexican dolphins; UD: Unknown dolphins. Bold numbers represent significant $p$-values; Bonferroni’s correction $p$-value = 0.0033. Values above the hyphen represent $p$-values; those values below the hyphen represent the genetic distances.

Given the bi and trimodal distributions described before, we performed a PCoA analysis using the putative origin as a criterion. We created a genetic structure using all individuals to delve into this possibility; the Bayesian method revealed two inferred genetic clusters ($K = 2$; $p = 1$; Figure 1).
Figure 1. Genetic structure of the *T. truncatus* populations analyzed in this study (four putative geographic populations and one population with unknown geography). (A) Bar plot of the several individuals analyzed; each vertical line (x-axis) is a single individual with colors representing each cluster’s membership proportion. Colors (red and green) represent the subpopulations conforming to the putative geographic origin. (B) Mean $L(K)\pm SD$ over five runs for each K value. Note: AD: Australian dolphins; CD: Cuban dolphins; JD: Japanese dolphins; MD: Mexican dolphins; UD: Unknown dolphins.

These data were used to check whether the putative geographic origin could suggest certain similarities with those inferred with structure. The MDS plot depicted similarities among MD, CD, and UD putative origin populations. MD2 and CD exhibited a marginal similarity ($p = 0.031$). Nevertheless, MD1 and MD2 did not differ significantly regarding UD (Figure 2). Likewise, JD presented a marginal difference with MD1. These data suggest that the putative MD1, MD2, CD, JD, and UD could form the first subpopulation. By contrast, AD was set apart from the rest of the groups.
4. Discussion

Genetic tools are indispensable for maintaining diversity with implications for reintroduction and conservation of endangered and vulnerable species. Statistical parameters and population genetic studies are crucial to validate the robustness of the markers selected; a critical point for their choice [40]. Particularly, the effectiveness and the informative degree of microsatellites used in the cetacean characterization have not been fully studied. Our particular interest was to know the effectiveness of genetic parameters and kinship relationships to guarantee biodiversity and implement identification strategies. The present study reports the genetic structure, the DNA fingerprint effectiveness, and the genetic diversity patterns of 210 *T. truncatus* individuals living captive from 18 Mexican centers using 15 STRs.

Previous studies have used multiple genetic markers for identification purposes [24–29]. Nonetheless, the sample sizes have been modest, whereas the statistical parameters have been restricted to the number of alleles, heterozygosity, and PIC. Such parameters are not informative and lack the effectiveness guideline, increasing the research costs. Knowledge of the statistical effectiveness parameters is crucial for several disciplines. Our study validated the statistical parameters of a set of 15 markers, reporting its specifications regarding kinship relationships and discrimination power. The set of markers proposed herein exhibited a remarkable discrimination capability ($1.141 \times 10^{-16}$), efficiently in distinguishing one individual in a range of millions of dolphins. Such effectiveness was
comparable with the data obtained from 19 STRs, although more cost-effective [22]. These features were the greatest strengths of our study, reinforcing its use in several disciplines. However, the prior studies on these microsatellites were essential for our study [24–29].

Of note, the probability of identity was obtained assuming random mating. Notably, the $F_{IS}$ values found suggest “inbreeding”, impacting the genetic variance and covariances between relatives [41]. Hence, this proficiency could be nuanced, depending on genetic closeness, and should be interpreted with caution [42]. Such “inbreeding” might reflect certain adaptations to captivity [43,44]. The human-induced selection over short time frames could also be causal to the loss of alleles and genotypes considered beneficial in wildlife [45]. Furthermore, using these 15 STRs could be extremely valuable in conservation and management strategies of endangered and vulnerable species, which requires several loci and molecular assessments to monitor the diversity [24]. Given that these STRs are highly conserved across odontocetes, their employment could encompass several genera, thereby adding to the scope of our findings. Several markers of the present study have been amplified in protecting species belonging to genera such as Balaena, Balaenoptera, Delphinapterus, Eschrichtius, Globicephala, Grampus, Megaptera, Stenella, Orcinus and Pontoporia, to mention a few [24,27,28,46].

Even though T. truncates is not considered an endangered species, these markers allowed us to determine its genetic structure by identifying two well-defined populations [8]. Such stratification could be associated with the capture habitat and possibly with the different ecotypes (i.e., inshore and offshore) [20,47–50]. The HWD and the remarkable LD found in the whole population are signatures of population stratification, confirmed through several analyses. Especially, the inshore ecotype has been reported in the Atlantic and Caribbean dolphins [47,49,51,52]. Coastal and offshore ecotypes have been documented in the Gulf of Mexico, being the inshore ecotype almost six times more prominent than the other one [49,53]. Genetic and isotopic differentiation have also been reported between the Gulf of California and the west coast of Baja California [54]. Thus, it is likely that some ecotype was shaping the genetic architecture of the samples, with the offshore one being the most probable given its high frequency [55,56]. Our data are discrepant from prior studies in other regions where low genetic diversity has been reported [57]. Such differences reinforce the necessity of genetic studies and a validated set of markers to expand the knowledge about diversity within Tursiops. These studies should also consider the environmental changes that have altered the genetic structure, suggesting that geography is only one of the different aspects contributing to genetic differentiation [16].

On the other hand, the knowledge of the genetic structure has been associated with the reintroduction facility, being successful where both populations (reintroduced and acceptor) share genetic similarities [17]. Diversity generation is also considered to minimize the inbreeding rates and ensure persistence [44,58]. In turn, the genetic structure is a cornerstone to determining survival and thriving [17]. Thus, before the reintroduction, random sampling from some wild individuals belonging to the putative acceptor populations should be performed to identify the genetic profiles [58]. In this setting, the proposed set of markers could be particularly valuable. Despite their utility, these 15 STRs should be accompanied by markers such as mitochondrial DNA (mtDNA) and other, more cost-effective markers such as the visual cues provoked by the damage to the tip of the dorsal fin [23,51]. Particularly, mtDNA is useful for identifying the migratory patterns and even identifying species and subspecies [22,59]. Although the markers employed herein exhibited a high identification ability, genome-wide genetic variation has been considered the best approach to maintaining the species’ biodiversity [60]. Nonetheless, this technology’s high costs are still prohibitive to developing countries such as Mexico. Alternatively, this set of markers, along with radio frequency identification and the permanent scars, might help avoid impersonation and illegal trafficking of these and other marine mammals.

Regarding the remarkable diversity found, we did not rule out that it could be related to other species such as Tursiops aduncus (T. aducus) that could have been confused with Tursiops truncatus truncatus (Tursiops t. truncatus). The genetic characterization using
STRs and mtDNA has shown the presence of *T. aducus* in coastal waters of southern Australia [61]. Unfortunately, the absence of non-nuclear genetic markers (i.e., control, D-loop, and Cytochrome regions) to clarify this doubt limited our study [22,49,62]. Another approach could be the analysis of genetic distances and their visualization in an MDS plot in Australian *Tursiops t. truncatus* and *T. aduncus*. Nonetheless, not all authors share their raw data or analyze the same loci, hindering such analysis. In this setting, the “inbreeding” exhibited by the studied populations was inconsistent with the highest diversity values and the greatest number of alleles, reinforcing this possibility.

Although genetic characterization is a key topic in the conservation and reintroduction process, it is only one of the diverse variables that should be considered [17,43,63]. Variables such as habitat, climate requirements, and management of pathogens carried by the reintroduced species, among others, must be considered in the reintroduction planning and individual management [63]. As regards the captive *T. truncatus*, their reintroduction is not feasible, mainly because they are not vulnerable populations and are dependent on humans. Foraging strategies and social relationships have been modified in captive dolphins due to human dependence [13]. In such circumstances, these cetaceans look forward to approaching vessels and people with subsequent implications such as injury from ingestion or even death [13]. Thus, reintroduction biology integrates multiple fields of study [58].

As mentioned before, the new regulations (i.e., article 60 Bis) did not prohibit scientific research and education programs [11] Thus, the collected information from ex situ dolphins could be extrapolated to wildlife specimens, promoting their welfare [8,9]. Research programs in high-income countries have provided substantial information about behavioral, cognitive, and physiological attributes, disease risk, and even the education of visitors, impacting their sensitivity and appreciation for these mammals [9,14,64,65]. Specifically, our markers could contribute positively to the characterization of biobanks and, subsequently, to reproductive assistance programs [9]. Hence, research programs represent a relevant opportunity to expand our knowledge about these and possibly other odontocetes [9,66].

5. Conclusions

In summary, the results presented herein validated using these 15 STRs to evaluate the probability of identity and to characterize the genetic structure with remarkable efficiency. These hypervariable markers are powerful and sensitive, enabling the genetic characterization of this and other species at a low cost, preserving the diversity amongst individuals. Nevertheless, more studies involving uniparental markers, sex determination, cranial and other morphological measurements should be determined to complete such a panorama.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ani12141857/s1, Table S1. Information about captive individuals and genotypes. Table S2. Allele frequency and statistical and paternity genetic parameters regarding the 15 loci studied in the four putative groups of *T. truncatus*. Table S3. Null allele frequencies of the 15 loci studied in the four groups of *T. truncatus*. Table S4. Linkage disequilibrium amongst all 15 loci studied in the four groups of *T. truncatus*. Figure S1. Map of the Mexican Republic showing the geographical area where the current captive centers (prior recreational centers) are found. The table exhibits the name of the political state with the acronym and the number of specimens, including only those dolphins with putative geographic assignments.

**Author Contributions:** Conceptualization: J.J.M., J.L.A.-F. and R.G. Methodology: R.M.N.-B., R.G., A.P.-M. and P.A.V.-D. Formal analysis: R.G., R.M.N.-B. and J.J.M. Funding acquisition: J.J.M. and J.L.A.-F. Analyzed data: R.G. and J.J.M. Resources: J.J.M. and J.L.A.-F. Writing—original draft preparation: R.G. and J.J.M. Writing—review and editing: R.G. and J.J.M. Supervision: R.G., J.J.M. and J.L.A.-F. All authors have read and agreed to the published version of the manuscript.

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