A New Dolphin Species, the Burrunan Dolphin *Tursiops australis* sp. nov., Endemic to Southern Australian Coastal Waters

Kate Charlton-Robb1,2*, Lisa-ann Gershwin3,4, Ross Thompson2, Jeremy Austin5,6, Kylie Owen2, Stephen McKechnie1

1 Centre for Environmental Stress and Adaptation Research, School of Biological Sciences, Monash University, Clayton, Victoria, Australia, 2 Australian Centre for Biodiversity, School of Biological Sciences, Monash University, Clayton, Victoria, Australia, 3 Tasmanian Museum and Art Gallery, Hobart, Tasmania, Australia, 4 South Australian Museum, Adelaide, South Australia, Australia, 5 Australian Centre for Ancient DNA, University of Adelaide, Adelaide, South Australia, Australia, 6 Sciences Department, Museum Victoria, Carlton Gardens, Melbourne, Victoria, Australia

**Abstract**

Small coastal dolphins endemic to south-eastern Australia have variously been assigned to described species *Tursiops truncatus*, *T. aduncus* or *T. maugeanus*; however the specific affinities of these animals is controversial and have recently been questioned. Historically ‘the southern Australian *Tursiops*’ was identified as unique and was formally named *Tursiops maugeanus* but was later synonymised with *T. truncatus*. Morphologically, these coastal dolphins share some characters with both aforementioned recognised *Tursiops* species, but they also possess unique characters not found in either. Recent mtDNA and microsatellite genetic evidence indicates deep evolutionary divergence between this dolphin and the two currently recognised *Tursiops* species. However, in accordance with the recommendations of the Workshop on Cetacean Systematics, and the Unified Species Concept the use of molecular evidence alone is inadequate for describing new species. Here we describe the macro-morphological, colouration and cranial characters of these animals, assess the available and new genetic data, and conclude that multiple lines of evidence clearly indicate a new species of dolphin. We demonstrate that the syntype material of *T. maugeanus* comprises two different species, one of which is the historical ‘southern form of *Tursiops*’ most similar to *T. truncatus*, and the other is representative of the new species and requires formal classification. These dolphins are here described as *Tursiops australis* sp. nov., with the common name of ‘Burrunan Dolphin’ following Australian aboriginal narrative. The recognition of *T. australis* sp. nov. is particularly significant given the endemism of this new species to a small geographic region of southern and south-eastern Australia, where only two small resident populations in close proximity to a major urban and agricultural centre are known, giving them a high conservation value and making them susceptible to numerous anthropogenic threats.

**Introduction**

Delphinids are the most ecologically diverse cetacean, occurring across a range of latitudes, in coastal and oceanic waters, and in estuarine and freshwater habitats [1]. In the last 25 years molecular techniques have markedly improved our understanding of cetacean taxonomy, including recognition of undescribed taxa within family Delphinidae [2,3]. However, relationships within sub-family Delphininae remain uncertain [4,5,6], largely due to their rapid global radiation and the ability of species to locally adapt [4]. Several species are distributed globally but show fine scale local population structure [7].

The genus *Tursiops* has been plagued with controversy with historically upwards of 20 species described, all synonymised with *T. truncatus* [8]. Only recently *T. aduncus* has been revalidated as the second *Tursiops* species, this based on morphological and mitochondrial DNA data [5,9,10,11]. In fact numerous studies have demonstrated that *Tursiops* is polyphyletic [5,12,13,14,15]. However, there is still controversy with two new distinct *Tursiops* species recently suggested [12,13,14]. In Australia, all *Tursiops* species have been historically recognised as *T. truncatus* [16]. However, Möller and Beheregaray [17] genetically confirmed the presence of *T. aduncus* off eastern Australia, while in Western Australia *aduncus* and *truncatus*-type haplotypes are also present [18].

In south-eastern Australia, morphological variation within *Tursiops* has been described for several decades [16,19,20]. In 1919, Scott and Lord [19] detailed the external and skeletal morphology of a unique, sexually dimorphic, southern form of *Tursiops* (as *T. tursio*). A single male specimen was captured by H.H. Scott in 1902 in the Cataract Gorge, Launceston, Tasmania. At the time, media reports, exhibition signage, and Scott’s own handwritten notes (held in the Queen Victoria Museum and Art Gallery) indicated that he believed it belonged to a distinct southern form of *T. tursio*. In 1914, a female specimen was obtained in the North Esk, Launceston, Tasmania. As stated, Scott...
and Lord [19] believed that the male and female belonged to the same species, and accounted for their numerous morphological differences by adding sexual dimorphism to the list of characters separating this southern form from the northern form of *Tursiops*. Iredale and Troughton [21] formally named Scott and Lord’s "holotype" has been listed as unknown [23]. We have recently located the male and female syntypes of *T. maugeanu* and incorporated them into this contemporary analysis.

In the current study, morphology indicates two forms of 'bottlenose' dolphin in south-eastern Australia, a physically smaller coastal form in semi-enclosed water bodies, and a larger more robust 'offshore' form. Locations of beach-cast dolphins suggest these two forms are parapatric, at least across some of their range. The smaller coastal form has been noted as both *T. truncatus* and *T. aduncus* [24,25] and due to the historical and current ambiguity of species identification, this form has more recently been referred to as *Tursiops* sp., southern Australian bottlenose dolphin (SABD) [13,14,26,27].

Charlton et al. [13] using the mtDNA control region first highlighted the divergent mtDNA lineage of SABD (using samples from the two Victorian populations), showing they did not cluster with *Tursiops*, *Delphinus* or *Stenella* species found world-wide. The average sequences divergence of the Victorian SABD to *T. truncatus* and *T. aduncus* (5.5% and 9.1% respectively [19]) was greater than that observed between recognised species within each of the *Cephalorhynchus* (2.5–4%) and *Lagenodelphis* (4.5–6.4%) genera [28]. Charlton et al. [13] concluding that these populations may represent an undescribed taxon. Möller et al. [14] later provided evidence for three genealogically distinct, reciprocally monophyletic, mtDNA lineages among the dolphins in southern Australia. Complementary microsatellite data indicated reproductive isolation among lineages [14]. Two of these lineages corresponded to published sequences of *T. truncatus* and *T. aduncus* [14]. The third lineage, including all SABD animals, was novel, the data suggesting it is a sister taxon of *Lagenodelphis hosei* (Fraser’s Dolphin). Kingston et al. [4] using mtDNA control region haplotypes from Charlton et al. [13] confirmed SABD as a monophyletic clade separate from *Tursiops* species, but with the sister taxa *Sousa chinensis* (Indo-Pacific Humpback Dolphin). Unlike other recently recognised species, Orcella heinsohni [3] and *Sotalia guianaensis* [2], where the classification within pre-existing genera was clear, SABD does not associate unambiguously with any described genus, and particularly not with either recognised species of ‘bottlenose dolphin’ in the genus *Tursiops*.

In 2004 a specialized Workshop on Cetacean Systematics was held to review cetacean taxonomy and provided criteria for species delimitation [29]. At that workshop it was agreed that multiple lines of evidence are required to demonstrate ‘irreversible divergence’ with criteria from both morphological and genetic data taken as proxies for reproductive isolation. The “ideal data set” will include both morphological data and data from multiple genetic loci [29].

Currently there are numerous Species Concepts, each with underlying properties that represent thresholds crossed by diverging lineages, different subgroups of biologist advocate different species concepts but they all exhibit underlying conceptual unity [30]. De Queiroz [30] highlights that unity and proposes a Unified Species Concept, stating that species are separately evolving metapopulation lineages on different evolutionary trajectories, the farther along process of divergence, the larger the number of differences. In this Unified Species Concept, any property (line of evidence) that provides evidence of lineage separation is relevant in species delimitations including genetic, morphological, ecological or behavioural [30]. A highly corroborated hypothesis of the existence of a new species requires multiple lines of evidence, the farther along the process of divergence, the easier it becomes to find and highlight evidence of separation. The Unified Species Concept [30] was used in the more recent 2009 Workshop for Defining Subspecies: Developing Guidelines for Marine Mammals [31] and whilst this Workshop was specific to the lower end continuum of subspecies differentiation, the Concept was used to highlight the “differentiation that characterizes the process of speciation” [31].

Since 2003 we have carried out extensive surveys, sampling and characterisation of ‘bottlenose’ dolphins from Victorian and Tasmanian coastal waters, using museum specimens, beach-cast strandings, live sightings and biopsies. In light of the confusion surrounding the taxonomy of these animals we use existing [13,14] and new genetic data, external and cranial morphometrics, incorporating the syntypes of *T. maugeanus*, and assess the taxonomic status of these animals. Consistent with the recommendations of the Workshop on Cetacean Systematics [29], and the Unified Species Concept [30] these multiple lines of evidence are used to establish the SABD as a new species of dolphin.

**Methods**

**Study location**

South-eastern Australia, encompassing coastal waters of Victoria and Tasmania (Figure S1). Southern Queensland, Australia (Museum specimens only).

**Cranial morphology**

Forty commonly used cranial measurements and tooth counts were taken from 44 specimens of ‘bottlenose’ dolphins from across Australia (Table 1 & 2; Table S1). Only adult specimens with complete data sets were used (those exhibiting secure fusion between maxillae and cranium [16]). All measurements were taken by the first author. Specimens were collected from locations across coastal Victoria (Museum Victoria (MV) (n = 26); and Monash University (MU) (n = 5) collections), Tasmania (Tasmanian Museum and Art Gallery (TMAG) (n = 5); Queen Victoria Museum and Art Gallery (QVMAG) (n = 5) collections) and Queensland Museum (QM) (n = 4) (Table 1). Cranial measures largely followed Kemper [20] and Wang et al. [32]. We included an undescribed measure, anterior pterygoid apex to palate (APAP); plus two undescribed qualitative features, shape of the palate and flattening on the maxilla at the base of the rostrum (Table S1).

As historically all *Tursiops* species were recognised as *T. truncatus*, QM specimens remain listed as either *Tursiops* sp. or *T. truncatus*. However *T. aduncus* is now known to be present in Queensland waters [33]. As such, we enlisted the technique used by Perrin et al. [34] in their assessment of the holotype specimen of *T. aduncus*, and used the range (min-max) of cranial measures presented in Wang et al [32], to conclude the four QM specimens were referable of *T. aduncus*.

**External morphology**

Eighteen external morphometrics (Table 3; Table S2) were taken from 17 ‘bottlenose’ dolphins from coastal Victoria (Table 1). Beach-cast dolphins were opportunistically measured during 2005–2009, by the first author and researchers at the Dolphin Research Institute, Monash University and the Department of Sustainability and Environment (Victorian Government). Animals
Table 1. List of all ‘bottlenose’ dolphin specimens examined with collection information.

| Museum - University code | Collection | MU Ext. Morph. | Date | Sex | Collection location |
|--------------------------|------------|----------------|------|-----|---------------------|
| MV | C29579 | 14/10/1985 | F | Western Beach | PPB |
| MV | C29587 | 1/06/1992 | M | Kennedy’s Point | WPB |
| MV | C29667 | 8/01/1987 | F | Ocean Grove | Vic |
| MV | C24944 | 2/06/1967 | F | Elwood | PPB |
| MV | C28760 | 13/11/1992 | - | Sandringham | PPB |
| MV | C29580 | 17/01/1986 | M | Murrells Beach | Vic |
| MV | C29577 | 23/07/1985 | F | Safety Beach | PPB |
| MV | C29586 | 27/07/1991 | M | Rippleside | PPB |
| MV | C10357 | - | - | - | - |
| MV | C31642 | - | - | - | - |
| MV | C35986 | 4/04/2006 | F | Mitchell River | Gips |
| MV | C35987 | 21/07/2006 | M | Hollands Landing | Gips |
| MV | C25071 | - | - | - | - |
| MV | Unknown | - | M | Murrells Beach | Vic |
| MV | C29506 | 16/04/1994 | F | Sorrento | PPB |
| MV | C11271 | - | - | - | - |
| QVMAG | 1365* | 11/11/1914 | F | North Esk River | Tas |
| TMAG | A1759 | 21/02/2003 | - | Marion Bay | Tas |
| TMAG | A2430 | - | - | - | - |
| QVMAG | 1946/7 | 16/01/1947 | M | North Esk River | Tas |
| QVMAG | 1972/1/35 | 1965 | M | Bass Strait | Tas |
| QVMAG | 1360** | 1902 | M | Cataract Gorge | Tas |
| MV | C31643 | - | F | - | - |
| TMAG | A2425 | - | - | - | - |
| TMAG | A198 | 1919 | - | East Coast | Tas |
| MV | C24987 | 18/05/1967 | Lorne | Vic |
| MV | C29585 | 13/05/1990 | M | Wild Dog Creek | Vic |
| MV | C29581 | 22/01/1986 | M | Port Fairy | Vic |
| TMAG | WAPSTRA | 2007 | - | - | - |
| QVMAG | MU270508 | 27/05/2008 | F | Cape Conran | Vic |
| MV | C35965 | 14/12/2006 | M | Lake Wellington | Gips |
| MV | C35985 | 1/12/2006 | M | Blonde Bay | Gips |
| MV | C35966 | 14/12/2006 | M | Lake Wellington | Gips |
| MV | C36750 | 4/11/2007 | M | Paynesville | Gips |
| MV | C35969 | 8/03/2006 | M | Phillip Island | Vic |
| MV | C35968 | 25/10/2007 | M | Tucker Point | Gips |
| MU | MU210108 | 21/01/2008 | M | Beaumaris | PPB |
| MU | MU23020308 | 23/01/2008 | M | Point Henry | PPB |
| MU | MU230607 | 23/06/2007 | F | Point Ricardo | Vic |
| MU | MU220108 | 22/01/2008 | F | Killarmy | Vic |
| MU | MU220108 | 22/01/2008 | F | Killarmy | Vic |
| MU | MU21108 | 2/11/2008 | M | Swan Reach | Gips |
| MU | MU291007 | 29/10/2007 | F | Jones Bay | Gips |
| MU | MU21007 | 23/04/2007 | M | San Remo | WPB |
| MU | MU190905 | 19/09/2005 | F | Corio Bay | PPB |
| MU | MU271006 | 27/10/2006 | M | Port Fairy | Vic |
| MU | MU280405 | 28/04/2005 | F | Kennett River | Vic |
| MU | MU010709 | 1/07/2009 | M | Portland | Vic |
were excluded from analysis if data was incomplete, bloating due to decomposition had occurred, or if the animal was a juvenile (less than 220 cm in length).

Morphological data analysis

Multivariate analyses of variance was used to test for sexual dimorphism in both cranial and external morphometrics datasets (not found, therefore males and females were pooled in further analyses) (cranial MANOVA, F₉,₁₉ = 1.289, p = 0.28, male = 17 female = 9; external MANOVA, F₁₁,₁₆ = 1.056, p = 0.449, male = 11 female = 6). Data were standardized by converting the raw data to z-scores. Hierarchical cluster analysis (HCA) using Euclidean pair-group average, discriminant function analyses (DFA) and principal component analysis (PCA) presented as supporting information only. DFA was also used to identify the measures which drove separation of haplotypes and if there was appropriate assignment into species ‘type’. Components analyse ((PCA) presented as supporting information only). Multivariate analyses of variance was used to test for sexual dimorphism in both cranial and external morphometrics datasets (not found, therefore males and females were pooled in further analyses) (cranial MANOVA, F₉,₁₉ = 1.289, p = 0.28, male = 17 female = 9; external MANOVA, F₁₁,₁₆ = 1.056, p = 0.449, male = 11 female = 6). Data were standardized by converting the raw data to z-scores. Hierarchical cluster analysis (HCA) using Euclidean pair-group average, discriminant function analyses (DFA) and principal component analysis (PCA) presented as supporting information only. DFA was also used to identify the measures which drove separation of clusters. MANOVA was used to test whether measures were statistically significant between clusters. All analyses were completed using SYSTAT v13 and PAST v1.94b.

Mitochondrial DNA sequencing

Skin samples were collected from beach-cast dead ‘bottlenose’ dolphins from coastal Victoria and stored in saline solution of 20% dimethyl-sulfoxide (DMSO), 0.25 M EDTA, saturated with NaCl, pH 7.5 [37]. Where skin samples were not available (museum specimens) tooth samples were collected. Numerous biopsy samples were taken from free-ranging dolphins during 2006–2008 using the PAXARMS biopsy system [38]. Several biopsy samples were used to verify genetic ‘type’ from the living populations (data not presented here).

Tooth samples were individually stored in sterilized Falcon tubes. Each tooth was sectioned and decontaminated by being submerged for 10 min in 12% sodium hypochlorite [39]. Sections were decalcified for up to four days using Morse’s Solution (10% Sodium Citrate, 20% Formic Acid) until ‘rubbery’ and flexible. Morse’s Solution was used as it does not degrade DNA quality [40]. Tooth samples were run from two separate extractions and two separate PCR reactions, including negative controls.

Total genomic DNA was extracted from skin and tooth samples using a Puregene Tissue kit (Gentra Systems) following manufacturer’s instructions, with modification for the teeth. Samples were extracted for analysis and quantity of genomic DNA using a NanoDrop ND-1000 Spectrophotometer. PCR amplification and sequencing of a ~450-bp fragment of the mtDNA control region and ~1200-bp of cytochrome b was undertaken following modified methods outlined in Charlton et al. [13] and Moller et al. [14] respectively. Tooth sample modifications include 1) an additional 4 µl of protease K during extraction (following manufacturer’s instructions), 2) use of 6 µl of 30 ng/µl gDNA in the PCR reaction due to reduction of quality and quantity of DNA compared to skin samples and 3) use of Bio-X-Act Short for all samples (Bioline). All PCR products were sent to Macrogen, Korea, for purification and sequencing. Capillary electrophoresis (CE) was conducted on an Applied Biosystems ABI 3730d DNA analyzer. Purified PCR products from tooth samples were also sequenced at Micromon, Monash University, with CE on an Applied Biosystems 3730S Genetic Analyser.

Syntype specimens of Tursiops mauganus

Small bone samples were taken from both syntype specimens of Tursiops mauganus [19,21] QVMAG#1365 and QVMAG #1360 using pre-sterilised 5 mm drill bits and a slow-speed hand drill. All PCR work was conducted at a dedicated ancient DNA facility (Australian Centre for Ancient DNA, University of Adelaide, South Australia) using stringent ancient DNA precautions and controls [41]. DNA was extracted from 100 mg bone powder using a modified Qiagen DNeasy Blood and Tissue kit [42]. A negative extraction control was included to monitor contamination during the extraction process. PCR amplification and sequencing of a 1124-bp region of the cytochrome b gene and 417-bp region of the control region was carried out using seven and three sets of primers respectively, each amplified a 132–200-bp overlapping fragment (Table S3). PCR amplifications were carried out in 25 µl reaction volumes containing, 2 mM MgSO₄, 0.25 mM each dNTP, 1 x PCR buffer (Invitrogen), 0.4 uM of each primer, 1 mg/ml RSA (Sigma) and 0.5 U of Platinum Taq HiFidelity (Invitrogen). PCR amplification was performed under the following conditions: 94°C 1 min, then 50 cycles of 94°C 15 s; annealing 55°C 15 s; 68°C 30 s, followed by a final elongation step of 68°C for 10 min. A PCR negative control and negative extraction control were included in all amplification attempts. PCR products were purified with Ampure (Agencourt) according to manufacturer’s instruction and Sanger sequencing was undertaken using the ABI prism Big Dye Terminator Cycle sequencing kit (PEApplied Biosystems, Foster City, CA). CE was carried out on an ABI 3130XL DNA analyser and raw sequences were edited using Sequencher (GeneCodes). To ensure authenticity and reliability of the sequence obtained from the historical specimens, all PCR and sequencing was repeated providing independent and duplicate coverage of all sequenced bases.

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Table 1. Cont.

| Museum - Collection | MU Ext. Morph. | Date | Sex | Collection location |
|--------------------|---------------|------|-----|---------------------|
| University code    | Collection    | Label |     |                     |
| JM1230             | QM            | -    | 6/02/1976 | - Moreton Bay | Qld |
| JM11375            | QM            | -    | 4/03/1996 | M Bargara Beach | Qld |
| 5241               | QM            | -    | 1983  | - Nth Stradbroke Is | Qld |
| 6428               | QM            | -    | 22/02/1987 | M Yellow Patch | Qld |
| 4155               | QM            | -    | -    | - Townsville       | Qld |

MV Museum Victoria; MU Monash University; QVMAG Queen Victoria Museum and Art Gallery; TMAG Tasmania Museum and Art Gallery; QM Queensland Museum; Tas Tasmanian waters; Vic Victorian coastal water; PPB Port Phillip Bay, Victoria; WPB Westernport Bay, Victoria; Gips Gippsland Lakes, Victoria; QLD Queensland waters.

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| Measure | Tursiops australis | Tursiops truncatus | Tursiops aduncus |
|---------|-------------------|-------------------|-----------------|
| n       | Mean (mm) | Range (mm) | n        | Mean (mm) | Range (mm) | n | Mean (mm) | Range (mm) | significance |
| min    | max      |           | min    | max      |           | min    | max      |           |             |
| BL     | 23      | 35.63     | 8      | 37.41    | 38.91    | 2      | 33.70    | 32.64     | 34.77       | ns           |
| CBL    | 27      | 493.58    | 13     | 527.88   | 505.5    | 5      | 441.00   | 424       | 455         | **           |
| DFWM   | 27      | 18.22     | 13     | 16.10    | 8.22     | 5      | 19.44    | 17.61     | 21.96       | ns           |
| DFWM   | 27      | 12.50     | 13     | 16.02    | 7.2      | 5      | 12.58    | 9.71      | 16.22       | *            |
| GLPT   | 27      | 60.12     | 13     | 75.89    | 67.16    | 3      | 63.71    | 61.49     | 66.88       | **           |
| GLPTF  | 27      | 114.59    | 13     | 115.85   | 108.8    | 5      | 193.90   | 180       | 200.50      | **           |
| GWPTF  | 27      | 213.35    | 13     | 237.62   | 224      | 5      | 219.90   | 213.50     | 227         | **           |
| GWEN   | 27      | 91.78     | 13     | 97.48    | 91.56    | 5      | 80.25    | 77.75     | 83.48       | **           |
| GWIN   | 27      | 66.98     | 13     | 76.64    | 60.1     | 5      | 63.16    | 56.85     | 68.73       | *            |
| GPRW   | 27      | 232.87    | 13     | 247.83   | 229.5    | 5      | 219.90   | 213.50     | 227         | **           |
| GPOW   | 27      | 162.64    | 13     | 168.62   | 135.78   | 5      | 196.87   | 190       | 200.50      | **           |
| GWPX   | 27      | 49.72     | 13     | 97.98    | 110.62   | 5      | 77.46    | 73.03     | 83.13       | ns           |
| GPARW  | 27      | 185.45    | 13     | 190.16   | 191.36   | -     | -       | -         | -           | -            |
| LAL    | 27      | 52.03     | 13     | 62.23    | 70       | 5      | 42.68    | 39.48     | 46.45       | **           |
| LO     | 27      | 69.95     | 13     | 69.06    | 71.16    | 5      | 60.63    | 54.60     | 64.80       | ns           |
| LTRL   | 27      | 232.87    | 13     | 247.83   | 229.5    | 5      | 219.90   | 213.50     | 227         | **           |
| LWM    | 27      | 142.19    | 13     | 149.54   | 163.94   | 5      | 119.90   | 109.79    | 127.41      | *            |
| ML     | 27      | 91.78     | 13     | 97.48    | 91.56    | 5      | 80.25    | 77.75     | 83.48       | **           |
| MFL    | 27      | 66.65     | 13     | 73.02    | 87.76    | 5      | 63.16    | 56.85     | 68.73       | *            |
| POL    | 22      | 34.20     | 8      | 34.04    | 31.8     | 2      | 30.83    | 29.90     | 31.77       | ns           |
| PRW    | 27      | 48.49     | 8      | 50.20    | 58.23    | 3      | 33.31    | 30.93     | 35         | ns           |
| RL     | 27      | 280.37    | 13     | 303.69   | 291.5    | 5      | 254.30   | 243       | 264         | **           |
| RWB    | 27      | 132.58    | 13     | 143.05   | 136.29   | 5      | 103.38   | 93.78     | 107.78      | **           |
| RW60   | 27      | 93.58     | 13     | 106.46   | 97.88    | 5      | 77.90    | 71.99     | 82.72       | **           |
| RWM    | 27      | 79.44     | 13     | 88.84    | 79.24    | 5      | 62.64    | 60.09     | 65.44       | **           |
| RW75   | 27      | 63.48     | 13     | 70.97    | 57.67    | 5      | 50.05    | 46.82     | 53.76       | **           |
| TREN   | 27      | 327.98    | 13     | 353.23   | 375.5    | 5      | 295.20   | 290       | 305         | **           |
| TRIN   | 27      | 333.43    | 13     | 360.58   | 339      | 3      | 297.17   | 292.50    | 305         | **           |
| UTLTR  | 27      | 236.59    | 13     | 253.04   | 240      | 5      | 209.10   | 200       | 216         | **           |
| VW     | 27      | 43.15     | 13     | 49.96    | 53.98    | 5      | 30.52    | 23.05     | 36.22       | **           |
| ZW     | 27      | 228.52    | 13     | 263.27   | 246      | 5      | 204.50   | 191       | 221.50      | **           |
| APAP   | 27      | 55.98     | 13     | 63.92    | 71.71    | 5      | 41.42    | 31.36     | 47.13       | **           |
| TPC    | 27      | 161.01    | 12     | 176.74   | 147.05   | 5      | 159.02   | 151.84    | 169.63      | **           |
| WAS    | 27      | 73.41     | 13     | 85.48    | 79.24    | 3      | 62.42    | 55.85     | 67.07       | **           |

**Tooth counts**

| Tooth counts | TLLL | TLLR | TTLL | TTUR |
|--------------|------|------|------|------|
| 25           | 22.84 | 21 | 26 | 28 |
| 25           | 23.12 | 21 | 26 | 28 |
| 25           | 23.88 | 22 | 27 | 28 |
| 25           | 23.85 | 22 | 28 | 28 |

*significant (p<0.05); **highly significant (p<0.001); ns not significant.
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Mitochondrial DNA sequence data analysis

All sequences were edited and aligned by eye using MEGA5 [43]. Each individual sequence was assigned to a haplotype. These haplotypes were used to confirm the genetic identity of the specimens represented in the morphological data and were also compared to the available sequences of both *Tursiops truncatus* and *T. aduncus*.

Whilst the mtDNA analyses was conducted on both cytochrome *b* and control region in the current study, in order to assess the phylogenetic affinities of these animals, comparisons must be made with additional taxa in the subfamily Delphininae. In order to achieve this we have chosen to use the mtDNA control region in isolation, as conducting a consolidation analyses using both mtDNA regions of the wider Delphininae taxa would involve taking individual haplotypes from GenBank and assuming individual identity and locale, and thus may misrepresent the affinities of each taxa. Mitochondrial DNA control region sequences representing multiple genera within subfamily Delphininae were downloaded from Genbank, including those previously reported in Charlton et al. [13] as AustVic, representing additional Victorian SABD haplotypes, and the *T. aduncus* holotype sequence (Genbank accession #DQ517442; Museum accession #ZMB66400). A model and parameters for the phylogenetic reconstruction were determined empirically using likelihood via MEGA5 [43]. The Bayesian Information Criterion scores (BIC) and Akaike Information Criterion, corrected (AIC) indicated Hasegawa-Kishino-Yano model [44] plus Gamma, with alpha (gamma, *K* = 5) = 0.3183 distribution, was the most appropriate model given the above data. The dataset was analysed using Maximum Likelihood (ML), Neighbour-Joining (NJ) and Bayesian inference. The ML analysis was conducted in MEGA5 [43] and the NJ analysis was conducted in PAUP v4.0b10 [45] using the above model. Reliability of the nodes for all trees was assessed using 1,000 bootstrap replicates. Bayesian phylogenetic inference was conducted through MrBayes 3.1.2 [46]. The Monte Carlo Markov Chain (MCMC) was run over 10,000,000 iterations, with a sampling frequency of 1,000. All other parameters in MR Bayes were set to default. The analysis was run over 2 replicates to assure convergence on a topology. The *Lagenorhynchus acutus* sequence was used as outgroup (see Table S6 for Genbank accession numbers).

Animal Ethics and Research Permit approval

Collection of samples was approved by Monash University Biological Sciences Animal Ethics Committee (AEC approval BSCI# BSCI/2008/21) and Victorian State Government, Department of Sustainability and Environment (DSE) Wildlife Act 1975 Research Permit (Permit No: 10005013).

Results

Cranial morphology

Hierarchical cluster analysis was performed on all cranial variables for 44 specimens and showed three highly supported groups (cophenetic correlation of 0.8337) (Figure 1a). Group 1 was largely represented by specimens collected in enclosed coastal waters of Victoria, Group 2 was largely represented by specimens collected from ‘offshore’ coastal waters of Victoria and Tasmania, whilst Group 3 was represented only by Queensland specimens.
Figure 1. Graphic analyses on cranial morphology delineating *Tursiops australis* sp. nov. and *Tursiops* species (A–B). Red = Group 1: *Tursiops australis* sp. nov., blue = Group 2: *T. truncatus*, green = Group 3: *T. aduncus*. Individuals with known mtDNA sequence are indicated by with the appropriate species colour code. (A) Hierarchical multivariate cluster analysis on cranial morphological traits showing three highly supported groups (cophenetic correlation 0.8337). *Tursiops australis* sp. nov. holotype (QVMAG #1365) in Group 1, and *Tursiops maugensis* male (QVMAG #1360) in Group 2. (B) Discriminant function analyses scatterplot of canonical scores on cranial morphological traits delineating *Tursiops australis* sp. nov., *T. truncatus* and *T. aduncus*.

**Figure 2.** Graphic analyses on external morphology delineating *Tursiops australis* sp. nov. and *Tursiops truncatus* (A–B). Red = Group 1: *Tursiops australis* sp. nov., blue = Group 2: *T. truncatus*. Individuals with known mtDNA sequence are indicated by with the appropriate species colour code. (A) Hierarchical multivariate cluster analysis on external morphological traits showing two highly supported groups (cophenetic correlation of 0.747). (B) Discriminant function analyses on external morphological traits delineating *Tursiops australis* sp. nov. and *T. truncatus* (Hotellings t2: p = 0.0224).

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doi:10.1371/journal.pone.0024047.g002
Table 4. Mitochondrial DNA cytochrome b region diagnostic sites.

|               | 1 | 1 | 1 | 2 | 2 | 2 | 2 | 2 | 3 | 3 | 3 | 3 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 5 | 5 | 5 | 5 | 5 | 5 | 5 | 5 | 5 | 5 |
|---------------|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| Tursiops australis sp. nov. |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Burru Cytb1   | C | A | T | G | T | C | T | G | T | T | A | C | T | T | C | C | C | C | C | C | C | T | A | T | C | C | C | G | A | T |   |   |   |   |   |
| Burru Cytb3   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Burru Cytb4   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Tursiops truncatus |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| T. maugeanus   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TCytb5        |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TT Cytb5      |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TT Cytb12     |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TT Cytb14     |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Tursiops australis sp. nov. |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Burru Cytb1   | T | C | T | C | T | C | T | C | C | C | G | T | T | T | T | A | C | T | C | T | C | G | C | T | A | C | C | T |   |   |   |   |   |
| Burru Cytb3   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Burru Cytb4   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| Tursiops truncatus |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| T. maugeanus   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TCTCytb5      |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TT Cytb5      |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
| TT Cytb5      |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
Discriminant function analysis (DFA) was used to determine whether cranial characteristics would distinguish the same groupings identified by the cluster analysis. The DFA scatterplot of canonical scores clearly show the three well separated ‘groups’ (Figure 1b). A posteriori classifications were 100% correct. Canonical Discriminant function weighting identified rostrum measures RWM and RW60, width measure GPOW (greatest postorbital width) and length measure TRIN (tip of the rostrum to internal nares) as important characters defining the groups (Table S7). Thirty-two of the 36 skull measures differed significantly between the ‘groups’ (Wilks’ λ MANOVA, $F_{66,18} = 12.839, p<0.001$) (Table 2).

External morphology

Hierarchical cluster analyses performed on 14 variables using 17 specimens clearly showed two highly supported groups (cophenetic correlation of 0.747) (Figure 2a).

DFA was also used on external characters to ascertain whether specimens were classified into the same groups as the cranial analyses. The histogram of dolphin specimens along the discriminant axis clearly show the two ‘groups’ well separated (Hotellings t2 $p = 0.022$) (Figure 2b). Again, a posteriori individual group assignments corresponded exactly. Discriminant function weighting showed several length measures (UJBH, UJEYE, UJGAP) and width of flukes (WFLU) as important characters defining the groups (Table S8). Nine measures of the 14 measures differed significantly between the ‘groups’ (Wilks’ λ MANOVA, $F_{11,2} = 64.32, p<0.001$) (Table 3).

Molecular analyses: Mitochondrial DNA sequence data

Molecular analyses were limited to samples included in the cranial (n = 18) and external (n = 17) morphology analyses from south-east Australian samples (ten of the animals were represented in both cranial and external datasets). DNA data could not be obtained from QM skulls. From the 25 samples where DNA was available no T. aduncus mtDNA types were found.

A 1086-base sequence of the mtDNA cytochrome b region was obtained from 18 samples representing both cranial and external morphology groups (Table S9). Six unique haplotypes were identified, three representing T. australis sp. nov. and three representing T. truncatus, defined by 62 variable sites (59 transition substitutions, 3 transversion substitutions). Forty-eight fixed site differences were noted between the two species (Table 4). Tursiops australis sp. nov. showed minimal intra-specific variation, with three variable sites, whilst T. truncatus showed more variation with 13 variable sites. Genetic sequences from the current study have been deposited on GenBank (Table S10).

Cytochrome b

A 1086-base sequence of the mtDNA cytochrome b region was obtained from 18 samples representing both cranial and external morphology groups (Table S9). Six unique haplotypes were identified, three representing T. australis sp. nov. and three representing T. truncatus, defined by 62 variable sites (59 transition substitutions, 3 transversion substitutions). Forty-eight fixed site differences were noted between the two species (Table 4). Tursiops australis sp. nov. showed minimal intra-specific variation, with three variable sites, whilst T. truncatus showed more variation with 12 variable sites.

Control region

A 418-base sequence of the mtDNA control region was obtained from 21 samples representing both cranial and external morphology groups (Table S9). Eight unique haplotypes were identified, three representing T. australis sp. nov. (two of which have previously been reported [13]) and five representing T. truncatus, defined by 30 variable sites (25 transition substitutions, five transversion substitutions and one single based insertion/deletion), when also including haplotypes previously reported in Charlton et al [13]. Ten diagnosable fixed base pair differences were found between the species (Table 5). In a similar way to the cytochrome b region, T. australis sp. nov. showed less intra-specific variation (three variable sites) when compared to T. truncatus (13 variable sites). Genetic sequences from the current study have been deposited on GenBank (Table S10).

| Table 4. Cont. |
|----------------|
| Discriminated sites separating south-east Australian Tursiops australis sp. nov. (Burru) and T. truncatus (TT) for mtDNA-CYTO b DNA sequence. These discriminate the diagnostic charaters between the two species (Table 4). Tursiops australis sp. nov. showed minimal intra-specific variation, with three variable sites, whilst T. truncatus showed more variation with 12 variable sites. Genetic sequences from the current study have been deposited on GenBank (Table S10). |

| Diagnostic sites separating south-east Australian Tursiops australis sp. nov. (Burru) and T. truncatus (TT) for mtDNA-CYTO b DNA sequence. These discriminate the diagnostic charaters between the two species (Table 4). Tursiops australis sp. nov. showed minimal intra-specific variation, with three variable sites, whilst T. truncatus showed more variation with 12 variable sites. Genetic sequences from the current study have been deposited on GenBank (Table S10). |
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| Diagnostic sites separating south-east Australian Tursiops australis sp. nov. (Burru) and T. truncatus (TT) for mtDNA-CYTO b DNA sequence. These discriminate the diagnostic charaters between the two species (Table 4). Tursiops australis sp. nov. showed minimal intra-specific variation, with three variable sites, whilst T. truncatus showed more variation with 12 variable sites. Genetic sequences from the current study have been deposited on GenBank (Table S10). |
Concordance between morphological and molecular groups

In order to assess which species the morphological ‘groups’ genetically represented, the individual’s mitochondrial DNA haplotype were overlaid on both cranial and external morphological datasets (samples highlighted by ★ in Figure 1a & 2a). The two distinct ‘groups’ from south-eastern Australian specimens concurred perfectly with Tursiops australis sp. nov. (Group 1) and southern form T. truncatus (Group 2) (Figure 1 & 2).

Tursiops maugeanus specimens

As stated, cranial and external morphology analyses presented several distinct groups. In all cases Group 1 incorporated the female T. maugeanus specimen (QVMAG #1365) and Group 2 incorporating the male T. maugeanus specimen (QVMAG #1360) (Figure 1a). MtDNA sequences (cytochrome b and control region) place the female T. maugeanus (QVMAG 1365) within Tursiops australis sp. nov. and the male T. maugeanus holotype (QVMAG 1360) within T. truncatus (Table 4 and 5; Figure 3).

Phylogenetic analyses

Phylogenetic reconstructions by Maximum Parsimony (MP), Neighbour-joining (NJ) and Bayesian analyses showed Tursiops australis sp. nov. clearly distinct from both Tursiops species, and in a monophyletic clade outside of any reported genera (Figure 3a & b). MP and NJ analysis methods showed very similar topologies, with minor discrepancies overall. As such we present here a consensus tree of the ML and NJ analysis of the mtDNA control region (Figure 3a). The tree was characterised by low level of resolution for most nodes, however bootstrap support for differentiation of each species was more robust (Figure 3a). Bayesian inference analysis showed only one slight variation in topology with the placement of Sousa chinensis (Figure 3b). Both ML and NJ phylogenetic reconstruction show a sister relationship to Stenella longirostris, whilst Bayesian analyses showed a sisters relationship to S. longirostris and also Lagenodelphis hosei.

Nomenclatural Acts

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TAXONOMIC TREATMENT. In our quantitative and qualitative morphological and molecular comparisons of the designated southern form T. maugeanus syntypes together with specimens from numerous strandings over the past century, it is clear that the two syntype specimens of T. maugeanus comprise two different species (Figure 1a; Table 4 & 5). In all cases, morphological and molecular, the male (QVMAG 1360) concurred with the southern hemisphere T. truncatus, while the female (QVMAG 1365), concurred with the undescribed species, SABD.

To clarify the taxonomy of T. maugeanus we identified two alternative taxonomic treatments. One option was to designate the female as the lectotype of T. maugeanus and thus resurrect the name; with this option, the paralectotype male would simply be subsumed under T. truncatus. However, Scott’s handwriting on the specimen label of the male, as well as his own extensive published and unpublished notes (held at QVMAG), make it clear that the species he envisaged was based on the male. Therefore, and given the current uncertain state of relationships within Delphininae [4,5,6], if it is someday demonstrated that the southern form of T. truncatus sensu Scott and Lord [19] and Iredale and Troughton [21] is distinct from the northern form, and if the female retained the name, our action would leave the name T. maugeanus, in essence, assigned to the wrong form. The other option was to designate the male as the lectotype of T. maugeanus, and leave it as a questionable junior synonym of T. truncatus for now; with this option, the paralectotype female would then be left without an identity.

After considerable consultation, we are convinced that the most conservative and stable approach is the second option above, thus leaving T. maugeanus as the appropriate available name for the southern form of T. truncatus, should it be found distinct from the northern. We anticipate that this is likely to occur, given the historical conclusions [19,21] and in light of the recent designation of the T. aduncus holotype whereby the South African Indo-Pacific form would be the name bearer of T. aduncus, leaving the western Pacific/Southeast Asian form T. aduncus possibly requiring a new name [34]. This then leaves the female paralectotype specimen, and the species she represents, needing a formal identity and thus also becomes available to be the holotype of the new species.

REVISED TAXONOMY OF TURSIOPS MAUGEANUS as TURSIOPS. TRUNCATUS

Order Cetacea Brisson, 1762
Family Delphinidae Gray, 1821
Subfamily Delphininae sensu LeDuc, 1999
Genus *Tursiops* Gervais, 1855

*Tursiops maugeanus* Iredale and Troughton, 1934

Synonymy

*Tursiops tursio*, southern form male. – Scott and Lord 1919: 96, pl. XXIII–XXV [in part].

*Tursiops maugeanus* - Iredale and Troughton 1934: 68, nom. nov. for *T. tursio* (southern form) Scott and Lord 1919 [in part].

*Tursiops truncatus* - Möller et al. 2008: 676; Kemper 2004: 42; Perrin 2009.

Lectotype here designated

QVMAG 1360, Cataract Gorge, Launceston, Tasmania, Australia, 1902.

Paralectotype

QVMAG 1365, Hobblers Bridge, North Esk River 5 km upstream from Tamar River, 11 November 1914; misidentified and does not in fact belong to *T. maugeanus* (=southern form of *T. tursio*), but represents an entirely new form requiring a separate name.

Revised Diagnosis

Body large, robust (mean 2.95 m in length; range 2.83–3.02 m); with a short rostrum (mean 11.6 cm; range 11–12.5 cm); with tall and falcate dorsal fin (mean 29 cm in height; range 23.5–34 cm); with two-banded colouration dorsally slate grey-black, ventrally off-white; lacking a pale shoulder blaze and ventral spotting. Skull is large and robust (mean 527.88 mm; range 505.5–547 mm), the rostrum is short (mean 143.05 mm; range 136.29–158.89 mm) and wide across all measures (Table 2), with shape of the suture between the palatine and maxilla being shallow triangular or flattened (mean 42.18 mm; range 23.88–57.84 mm); ratio between the pterygoids and palatine is approximately 2:1; with obvious ‘pinched’ appearance where the maxilla transitions into the premaxilla (Figure 1). On average has 90 teeth (22 lower left; 22 lower right; 23 upper left; 23 upper right).

Molecular diagnostic characters

See below comparison with *Tursiops australis* sp. nov.

Remarks

Concordant results from multiple independent data sets suggest that the syntype specimens of *Tursiops maugeanus* belong to two different species (Figure 1; Table 4 & 5). The male lectotype specimen of *T. maugeanus* is identical in its morphological and molecular features to the offshore southern form of *T. truncatus* (Figure 1; Table 4 & 5). Therefore, we provisionally regard *T. maugeanus* as a junior synonym of *T. truncatus* under the current.
The taxonomic system. If the southern form of *T. truncatus* (hereafter referred as s.f. *T. truncatus*) is demonstrated to be a different species or subspecies from the northern form, the appropriate available name for the southern form would be *T. maugeanus*. The female paralectotype, which does not belong to *T. maugeanus*, is treated below.

**TAXONOMY OF NEW SPECIES.**

Order Cetacea Brisson, 1762
Family Delphinidae Gray, 1821

Subfamily Delphininae *sensu* LeDuc, 1999

Genus *Tursiops* Gervais, 1855

*Tursiops australis* sp. nov.

urn:lsid:zoobank.org:act:54BA663A-BDE6-4E12-A9D2-84F6793EF4EA

Figure 4 & 5

**Synonomy.**

*Tursiops turio*, southern form; female. - Scott and Lord 1919: 96, pl. XXIII–XXV [in part].

*Tursiops maugeanus* - Iredale and Troughton 1937: 68, nom. nov. for *T. turio* (southern form) Scott and Lord 1919 [in part].

*Tursiops sp.* - Scarpaci et al. 2003: 342; Warren-Smith and Dunn 2006: 357.

Victorian coastal bottlenose dolphin - Charlton et al. 2006: 173.

Southern Australian Bottlenose Dolphin - Möller et al. 2008: 676; Owen et al. 2011.

South Australian *T. truncatus* - Kingston et al. 2009:4.

*Tursiops truncatus* - Ross and Cockcroft 1990: 124.

*Tursiops aduncus* – Kemper 2004: 42.

Not *Tursiops truncatus* (Montagu, 1821): 75, pl. III.

Not *Tursiops turio* (Fabricius, 1780): 49.

Not *Tursiops aduncus* (Ehrenberg, 1832)

**Etymology**

Species name, *australis*, is in reference to the species link with Australia and is Latin for ‘southern’.

**Holotype.**

QVMAG 1365, Hobblers Bridge, North Esk River 5 km upstream from Tamar River, 11 November 1914; previously published as the female of the southern form of *Tursiops turio* by Scott and Lord, 1919, later named *Tursiops maugeanus* Iredale and Troughton, 1934. Repository location: Queen Victoria Museum and Art Gallery, Launceston, Tasmania, Australia.

**Paratypes.**

Monash MU210108, Beaumaris, Port Philip Bay, VIC, 21 Jan 2008; male. Monash MU230108, Point Henry, Geelong, VIC, 23 Jan 2008; male. MV C29579, Western Beach, Geelong, VIC, 14 Oct 1985; female. MV C29587, Kennedy’s Point, Westernport Bay, VIC, 1 Jun 1992; male. MV C29667, Ocean Grove, VIC, 8 Jan 1987; female. MV C24944, Elwood, VIC, 2 Jun 1967; female. MV C28760, Sandringham, VIC, 13 Nov 1992; sex unknown. MV C29580, Murrells Beach, VIC, 17 Jan 1986; male. MV C29577, Safety Beach, VIC, 23 Jul 1985; female. MV C29586, Rippleside, VIC, 27 Jul 1991; male. MV C35986, Mitchell River, VIC, 4 Apr 2006; female. MV C35987, Hollands Landing, VIC, 21 Jul 2006. MV C35985, Lake Wellington, VIC, 14 Dec 2006; male. MV C35966, Paddy Bay, VIC, 30 Aug 2006; male. MV C35985, Blonde Bay, VIC, 1 Dec 2006; male. MV C35966, Lake Wellington, VIC, 14 Dec 2006; male. MV C36750, Paynesville, VIC, 4 Nov 2007; male. MV C29506, Sorrento, VIC, 16 Apr 1994; female. TMAG A1759, Marion Bay, TAS, 21 Feb 2003; unknown sex.

**Type Locality.**

North Esk River, 5 km upstream from Tamar River at Hobblers Bridge, Launceston, Tasmania (type locality), Port Phillip Bay and Gippsland Lakes, Victoria, Australia.

**Diagnosis.**

**External morphology.** *Tursiops australis* is smaller (mean 2.57 m in length; range 2.27–2.78 m) than s.f. *T. truncatus* (mean
2.95 m in length; range 2.83–3.02 m) (Table 3), larger than T. aduncus (mean 2.25 m in length; range 14.0–268 m [11]). Rostrum is smaller and ‘stubbier’ (mean 10.9 cm; range 9.4–12 cm) than T. aduncus (mean 13.4 cm; range 8.0–15.5 cm [11]), similar to s.f. T. truncatus (as above). Dorsal fin is falcate like T. truncatus, c.f. the small triangular fin of T. aduncus. Tursiops australis has a tri-banded colouration grading conspicuously as follows: dark bluish-gray dorsally and on the sides of the head and body; light gray along the midline, extending as a pale shoulder blaze on the flank below the small triangular fin of an elongated triangular shape, in contrast to s.f. T. aduncus.

Skull morphology

The skull is more ‘petite’ than s.f. T. truncatus. Average skull length (CBL) is 493.58 mm (range 470–513 mm) smaller than that of s.f. T. truncatus (as above) and larger than T. aduncus (mean 441 mm; range 424–455 mm). Across all measures the rostrum is wider and shorter than T. aduncus (Table 2). The palate is long (mean 55.90 mm; range 45.27–63.85 mm) and the shape of the suture between the palate and maxilla is an elongated triangular shape, in contrast to s.f. T. truncatus and T. aduncus shallow triangular or flattened shape (mean 42.18 mm; range 23.68–57.8 mm and mean 41.42; range 31.36–47.13 respectively) (Table 2 and Figure 4). The ratio between the pterygoids and palate observed in T. australis is approximately 1:1, c.f. 2:1 for s.f. T. truncatus. On average T. australis has 94 teeth (23 lower left; 23 lower right; 24 upper left; 24 upper right). Teeth are long and conical, with older Gippsland Lakes and Tasmanian animals exhibiting substantial wear in the front and back teeth. The maxilla is flattened and smoothly transitional into the premaxilla toward the base of the rostrum, lacking the obvious the ‘pinched’ appearance of s.f. T. truncatus (Figure 4).

Molecular diagnostic characters

Tursiops australis differs from s.f. T. truncatus significantly at 58 diagnosable fixed base pairs across two mtDNA gene regions, 48 fixed site differences in a 1086-base sequence of the mtDNA cytochrome b region (Table 4) and 10 differences along a 418-base sequence of the mtDNA control region (Table 5).

Common Name

We propose the common name ‘Burrunan Dolphin’ for Tursiops australis. ‘Burrunan’ is an Australian aboriginal name given to dolphins (used in the Boonwurrung, Woiwurrung and Taungurung languages) meaning “name of a large sea fish of the porpoise kind” [47]. One of the two only known resident populations of T. australis is in Port Phillip Bay where the Boonwurrung people have documented their existence for over 1000 years.

Distribution

South-eastern and southern Australian coastal waters, including Victoria, Tasmania and South Australia (Figure S1). Two known resident populations of T. australis occur in Victoria; Port Phillip Bay (est. 90 animals [26]) and the Gippsland Lakes [13] where we estimate ~50 animals. Tursiops australis haplotypes have also been documented from dolphins in eastern Tasmanian waters [14] and coastal regions of South Australia in the Spencer Gulf region and west to St. Francis Island [14,48]. No T. australis haplotypes have been reported north of the Victoria/New South Wales border, or west of St. Francis Island, South Australia.

Discussion

Here we present clear and consistent molecular and morphological differences thus demonstrating the existence of a new species of dolphin in south-eastern Australian waters.

Relationships of Tursiops australis with other taxa

Morphological analyses reveal the new species and the two recognised Tursiops species differ in quantitative and qualitative cranial characters and in external morphology. The combination of overall size of the adult body, rostrum length and width, tail and falcate dorsal fin, the distinctive tri-colouration patterning and the extension of the white ventrum extending over the eye in T. australis (Figure 5; Table 3) differ conspicuously from the two recognised Tursiops species in Australian waters.

Cranial comparisons between T. australis and T. truncatus from south-eastern Australia and T. aduncus from Queensland, Australia (current study) show significant differences across multiple measures (Table 2). The three species grouped separately using multiple forms of statistical analyses (Figure 1 and S2). Tursiops australis overall size and shape of the skull is somewhat intermediary between the two recognised Tursiops species, however there are only a few characters that overlap in their range (Table 2). Two particular qualitative cranial characters, the shape of the suture between the palate and maxilla (quantifiable by a ratio between the length of the pterygoids and palate), and the smooth transition between the maxilla and pre-maxilla region (Figure 4) are clearly diagnostic of T. australis. When comparing T. australis to T. aduncus there is also clear differences. Tursiops australis shows a longer and wider skull to T. aduncus holotype specimen [34] and to reported T. aduncus from both South African and Chinese water [32] (Table S1). In addition the T. australis rostrum is significantly narrower across all measures and has more teeth (Table 2 and Table S1).

Further, animals grouped by external and cranial morphometrics as either T. australis or s.f. T. truncatus were in every case identified to the same group determined using molecular analysis (Figure 1). Charlton et al. [13] found high mtDNA control region sequence divergence between the new species and Tursiops truncatus (5.5%) and between the new species and T. aduncus (9.1%). Using mtDNA cytochrome b, Möller et al. [14] reported between 5.5% and 7.7% divergence between the new species and T. truncatus. This is larger than between T. truncatus and T. aduncus (3.2%–5.8%) [14], and between several other delphinid species that are grouped in the same genus, such as between Lagenorhynchus obscurus and L. obliquidens (1.22%) [49], Delphinus delphis and D. capensis (1.09%) [50], and between the recently described Sotalia fluviatilis and S. guianensis (2.5%) [2], and Orcella heinsohni and O. brevirostris (5.9%) [3]. We show this divergence is supported by clear diagnostic fixed sequence differences between T. australis and s.f. T. truncatus (cytochrome b = 48 fixed differences; control region = 10 fixed differences; Table 4 and 5 respectively). In addition, Möller et al [14] examined the new species (designated mtDNA clusters) using multiple nuclear markers and found evidence for complete reproductive isolation of the new species to both T. truncatus and T. aduncus. This high level of genetic divergence, complete reproductive isolation and the ambiguity of placement within any recognised genera strongly indicate that these coastal dolphins are not simply ecotypes of either recognised Tursiops species but are in fact representative of a new species. Irreversible divergence and distinct evolutionary trajectory of T. australis from recognised Tursiops species appears indisputable based on these multiple non-overlapping data sets.
Polphyly of Tursiops

As previously discussed, numerous studies have demonstrated that *Tursiops* is polyphyletic [5,12,13,14,15]. When assessing the phylogenetic relationships of *T. aduncus*, using the mtDNA cytochrome *b* and the control region, the sister taxa most commonly suggested is *D. delphis* and *S. corniculata* [5,13,14], in addition Möller et al. [14] has also suggested *S. frontalis* and *S. flibustieri* are also sister taxa. However based on mtDNA control region Kingston et al. [51] does not show the sister relationship between *D. delphis* and *T. aduncus* and based on AFLP places *T. aduncus* with *L. hosei* (using Nei-Lei neighbour joining anlaysis). Regardless of the DNA region used and phylogenetic analysis performed *T. truncatus* forms a separate clade from *T. aduncus* [5,12,13,14,51].

In this study we have also shown that *Tursiops* in polyphyletic, with *T. aduncus*, *T. truncatus* and *T. australis* on three independent lineages. Using all three phylogenetic analyses, the placement of *T. australis* is outside of both *Tursiops* species, with a sister relationship to *S. longirostris* (using ML and NJ methods) and additionally *L. hosei* using Bayesian inferences. Möller et al. [14] using mtDNA cytochrome *b* suggests the same sister relationships.

Alternative taxonomies

Unlike other recently recognised species, *Orcella heinsohni* [3] and *Sotalia guianensis* [2] where the classification within pre-existing genera was clear, this species based on multiple molecular regions [4,13,14] does not associate unambiguously with any existing genus. Whilst, as the discussed, the genus *Tursiops* is currently accepted as polyphyletic, Kingston et al. [4] states there is no support for a close genetic relationship between the two recognised *Tursiops* species, despite the morphological similarities, and along with others, calls on a review of not only the genus *Tursiops* but of family Delphinidae [4,5,12]. Natoli et al. [12] also raising the issue of generic affinities of *Tursiops*, more specifically the South African *aduncus*-type with the reported closeness to *D. delphis*, however no attempt was made at resolving the generic affinities. Given this current state of taxonomic uncertainty we believe that the most conservative approach at this time is to classify the new species in genus *Tursiops*, pending revision. We further believe that once revision of the Delphinidae is conducted, it is likely that this new species will be shown to represent a unique genus; if that is the case, we believe the genus name *Tursodepis* would be appropriate (from the Latin ‘tursio’, meaning ‘porpoise’, and Greek ‘delphis’, meaning ‘dolphin’).

In contrast, a number of nuclear DNA regions were also investigated in this study (data not shown). They include intron regions; CHRNA (283 bp) and POLA (330 bp) [52] and anonymous nuclear regions; Del10 (346 bp), Del 12 (575 bp) and Del 16 (533 bp) [53] for 19 individuals from *T. australis*, *T. truncatus*, *T. aduncus* and *L. hosei* (with species identification based on mtDNA regions). Of the five intron regions, four suggested no differentiation between the four species however one region (Del12) showed consistent species specific differences, defined by 3 variable sites (all transition substitutions). A possible explanation for this lack of differentiation may be due to the slower evolving nuclear regions, the rapid radiation of the delphinids (as also highlighted also by the current confused state of many generic affinities of dolphins [5,12,15]) and thus the potential of recent shared ancestry of these species. Caballero et al. [15] found significantly less parsimonious informative characters at each of the nine intron regions in comparison to each of the mtDNA control region and cytochrome *b*. In addition, the small samples size, taxa examined and lack of available Delphinidae GenBank submitted intron sequences for comparison may also be the limiting factor for species differentiation in this case. Larger sample sizes and greater representation from multiple taxa across Delphinidae would clearly be required to investigate generic affinities further, however, the clear and consistent morphological and molecular differentiation presented in this paper clearly support species level distinction.

Additional evidence

An additional line of evidence for separation of *T. australis* and s.f. *T. truncatus* is provided in Owen et al. [27] using stable isotope signatures of both species. Owen et al. [27] indicated *T. australis* (noted as SABD) was distinct from s.f. *T. truncatus* (noted as common bottlenose dolphin CBD), with s.f. *T. truncatus* having significantly lower values for δ¹³C and δ¹⁵N compared to *T. australis*. They conclude this distinction of the stable isotope signatures between the two species strongly indicates they forage in different areas and are likely to feed on different prey, thus providing an additional line of evidence for the recognition of *T. australis*.

Conservation value

Whilst, as previously discussed, there is an urgent requirement to undertake a full review of the Delphinidae Family, this manuscript is an important step in this review. This new species has erroneously been ‘labelled’ *Tursiops truncatus* and *T. aduncus*, of which we have demonstrated with clear and consistent evidence that it is neither. In addition, these dolphins have been ‘living under the eye’ in a well populated urban environment, have been the focus of multiple researchers and due to the multi-disciplinary approach taken in the manuscript we have been able to formally identify this species, thus highlighting the ‘need’ for other such studies to not ‘look in isolation’ of one line of evidence but to use a multiple disciplinary approach to assess the level of divergence.

The formal recognition of this new species is of great importance to correctly manage and protect this species, and has significant bearing on the prioritization of conservation efforts. This is especially crucial given it’s endemism to a small region of the world, with only two small known resident populations and the proximity of those to major shipping ports, commercial and recreation fisheries, residential, industrial and agricultural stressors. Recognition of this new species opens the pathway that *T. australis* would qualify for listing as a threatened species under the Australian Environment Protection and Biodiversity Conservation Act 1999 (EPBC Act) thus allowing immediate and directed conservation effort for further protection.

Supporting Information

Figure S1 Sampling locations of dolphins across Australia (TIF)

Figure S2 Principal component analysis on cranial measures; scatter plot of Principal components 1 & 2 (TIF)

Figure S3 Principal component analysis on external morphology measures; scatter plot of Principal components 1 & 2 (TIF)

Table S1 Cranial measures (DOC)

Table S2 External morphology measures (DOC)

Table S3 Principal component analysis loadings of the first three Principal components (PC) on 34 cranial measures from 40 ‘bottlenose’ dolphin skulls (DOC)
Table S4 Principal component analysis loadings of the first three Principal components (PC) for 11 external measures from 17 'bottlenose' dolphins (DOC)

Table S5 PCR primers used to amplify mtDNA cytochrome b gene and control region from Tursiops truncatus (DOC)

Table S6 GenBank accession numbers and species information for samples incorporated to phylogenetic analyses (DOC)

Table S7 Discriminant function analysis loadings on 34 cranial measures from 40 'bottlenose' dolphin skulls (DOC)

Table S8 Discriminant function analysis loadings for 11 external measures from 17 'bottlenose' dolphins (DOC)

Table S9 Species classification overview based on different analyses and characters (DOC)

Table S10 GenBank accession numbers of the sequences from this study (DOC)

Table S11 Average cranial measures (mm) and tooth counts for Tursiops aduncus, Tursiops truncatus, Tursiops aduncus (current study) and from Tursiops aduncus holotype (Perrin et al. 2007) and Tursiops aduncus (Wang et al. 2000) (DOC)

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Author Contributions

Conceived and designed the experiments: KCR SWM. Performed the experiments: KCR JA LG KO. Analyzed the data: KCR. Contributed reagents/materials/analysis tools: SWM JA RT. Wrote the paper: KCR. Provided thorough review of manuscript: LG RT SWM JA KO.

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