Mitochondrial variation among Australian freshwater turtles (genus Myuchelys), with special reference to the Endangered M. bellii

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ABSTRACT: Identifying species and the relationships among them remains important for assessing biodiversity trends and is a critical focus for reversing global biodiversity loss. The saw-shelled turtles of Australia, in the genus Myuchelys, show cryptic diversity and include species that range from Endangered (M. bellii), through those that are locally abundant but extremely limited in distribution (M. georgesi and M. purvisi) to those that are common and widespread (M. latisternum). The Endangered M. bellii is restricted to 3 small isolated populations in the headwaters of the Murray-Darling basin, in the Border, Gwydir and Namoi tributaries. There is no evidence of strong differentiation among these 3 populations based on mitochondrial DNA (mtDNA) divergences; rather there is only a shallow genetic structure ranging from 0.1 to 0.3% divergence. The 3 restricted and small populations of the Endangered M. bellii face a number of threatening processes and require conservation management across state boundaries as a single biological species. The mtDNA phylogeny supports previous phylogenetic findings of a deep phylogenetic divergence between M. purvisi and M. georgesi (13.5% mtDNA) and the sister taxa relationship of M. latisternum and M. bellii. A notable exception was our finding that Emydura macquarii and M. georgesi are sister taxa inside the Myuchelys radiation.

KEY WORDS: Emydura macquarii · Myuchelys georgesi · Elseya · Chelidae · Freshwater turtles · Cryptic species

INTRODUCTION

Biodiversity loss is of increasing concern to governments nationally and internationally. A substantial array of legislative and regulatory systems are in place to protect it, recognising the value the community places on biodiversity for the services it delivers in maintaining the quality of the air we breathe, the water we drink and the food we eat (Cairns 1997, Salzman 1998, Baumgärtner 2007, Anderson et al. 2009). While these legislative instruments recognise biodiversity at a range of levels, the central focus of biological conservation is at the species level, so identifying species, and relationships among species, remains a core focus of managers aiming to preserve biodiversity. For example, international systems of biodiversity protection such as the International Union for Conservation of Nature (IUCN) Red List and national systems such as the United Kingdom Endangered Species Act, United States Endangered Species Act and the Australian Environment Protection and Biodiversity Conservation Act (EPBC Act) rely on good taxonomic knowledge at the species level. However, because such knowledge is often incomplete, conservation action can be misguided or misdirected (Roman et al. 1999, Engstrom et al. 2004, Georges et al. 2011).

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Recent developments in molecular approaches to describe species genetic makeup have accelerated progress in documenting biodiversity and have revealed an unexpected level of cryptic biodiversity. For example, the Asian leaf turtle *Cyclemys dentata* was regarded as a single widespread polytypic species until molecular and morphological analysis revealed a species complex of several formerly cryptic species (Fritz et al. 2008, Stuart & Fritz 2008). The Endangered giant narrow-headed softshell turtle *Chitra indica* is also likely to be a species complex, as mitochondrial NADH dehydrogenase subunit 4 sequences reveal 3 deeply divergent, monophyletic lineages (Engstrom et al. 2002). Similarly, the European pond turtle *Emys obicularis* contains a cryptic species endemic to Sicily (Fritz et al. 2005, 2006). The Sicilian form, *E. trinacris*, is highly distinct genetically while morphologically resembling *E. obicularis galloitalica* (Fritz et al. 2005).

Failure to recognise underlying cryptic diversity at or below the level of species can have a great influence on biological conservation outcomes, either through declines in cryptic species that go undetected and as a result are poorly managed (as with *Cyclemys*, Fritz et al. 2008, Stuart & Fritz 2008) or because diversity that is suspected but with poor foundation leads to misallocation of resources (Georges et al. 2011). In addition, phylogenetic information has a role to play in assessing distinctiveness, an attribute that has an influence on determining conservation priorities (Faith 1992, Moritz 1995).

The saw-shelled turtles of Australia, in the genus *Myuchelys*, show cryptic diversity where species are morphologically conserved including suspected taxa within currently accepted species. The genus *Myuchelys* (formerly *Elseya latisternum* clade) was recently separated from the genus *Elseya* to account for the paraphyletic arrangement with respect to *Emydura* (Thomson & Georges 2009, Georges & Thomson 2010). The genus includes species that range from Endangered (*M. bellii*), through to those that are locally abundant but extremely limited in distribution (*M. georgesi* and *M. purvisi*) to those that are common and widespread (*M. latisternum*) (Fig. 1). *M. bellii* is of particular interest because it is restricted to 3 small isolated populations in the headwaters of the Murray-Darling basin, in the Border, Gwydir and Namoi tributaries. As such, the species is listed as Endangered in the IUCN Red Data Book (IUCN 2011) and vulnerable by the Australian authorities at State (Threatened Species Conservation Act, New South Wales 1995) and national level (EPBC Act, Commonwealth of Australia 1999). Suggestions that the populations in Girraween National Park of Queensland are sufficiently distinct to warrant recognition at species or subspecies level (Cann 1998, Wells 2002) is of particular interest to conservation agencies. (Note that Wells 2002 is not regarded as a publication for nomenclatural purposes. Refer to Georges & Thomson 2010.)

Multilocus allozyme studies (Georges & Adams 1992) revealed that *Myuchelys georgesi* and *M. purvisi* were deeply divergent genetically, although remarkably similar morphologically differing only in the presence or absence of neural bones in the carapace (Thomson & Georges 1996). They were regarded as the same taxon until *M. georgesi* was formally described by Cann (1997). Furthermore, they are not sister taxa, as the accompanying phylogenetic analysis by Georges & Adams (1992, 1996) revealed that their common ancestor had *M. latisternum* and *M. bellii* among its descendants. Recent taxonomic revisions (Thomson & Georges 2009, Georges & Thomson 2010) based on the earlier phylogenetic analysis place *Emydura* between *Elseya* and *Myuchelys*. These relationships have often been

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![Fig. 1. *Myuchelys* spp. Capture localities of the *Myuchelys* specimens used for mtDNA sequencing. Macquarie Marshes and Girraween National Park (Bald Rock Creek) are also shown. Drainage basin names are those of Australia’s River Basins 1997 available from Geoscience Australia. Inset depicts the distribution of *M. latisternum* from coastal catchments of northern New South Wales, eastern Queensland and the Northern Territory of Australia.](image-url)
contested in past taxonomic studies using morphology or genetic markers (Gaffney 1977, Legler & Cann 1980, McDowell 1983, Thomson & Georges 2009). Notwithstanding limitations of single locus markers, such as mtDNA, in species boundary studies, we examined mtDNA variation across the genus *Myuchelys* to determine whether the signal there is consistent with that of allozyme electrophoresis, particularly with regard to the relationship between the cryptic species *M. georgesi* and *M. purvisi* and to test the currently accepted phylogenetic hypothesis of monophyly of the genus with respect to *Enyдуra* and *Elseya* species. We also examined levels of mitochondrial divergence among 3 populations of the Endangered *M. bellii* to assess whether there is corroborating evidence in support of earlier suggestions that these be distinguished as separate species or subspecies. This study will assist in setting conservation priorities for *M. bellii*.

**MATERIALS AND METHODS**

Specimens of turtles within the genus *Myuchelys* (sensu Thomson & Georges 2009) were collected from within their natural ranges by a combination of baited traps (after Legler 1960) or by hand with the aid of mask and fins. The focal species, *M. bellii* (Gray 1844), was collected from the headwaters of the Namoi and Gwydir Rivers of New South Wales (NSW) and Border Rivers of Queensland (Qld), each draining into the Darling River of the Murray-Darling drainage basin. Other species were *M. purvisi* (Wells & Wellington 1985) restricted to the Manning River of coastal NSW, *M. georgesi* (Cann 1997) restricted to the Bellinger River of coastal NSW and *M. latisternum* (Gray 1867) from the rivers draining northeast NSW. Three species were chosen *a priori* as a nested series of outgroup taxa: the distantly related long-necked species *Chelodina rugosa* (Ogilby 1890) from rivers draining the Gulf of Carpentaria (Qld), *Elseya dentata* (Gray 1863) from the Roper River in the Northern Territory (NT), a close relative to *Myuchelys* genus, and *Enyдуra macquarrii* (Gray 1830, 1857) from the Murray River of Victoria (Vic), a species currently positioned taxonomically between *Elseya* and *Myuchelys* (Thomson & Georges 2009).

Skin samples (2–5 mm²) were taken from the neck or hind leg in the case of *Myuchelys bellii* using a sharp sterilised scalpel or surgical scissors. The samples were preserved in 95% ethanol and transported to the University of Canberra where they were stored at −20°C. Tissue samples for other species were obtained from the wildlife tissue collection of the Institute for Applied Ecology at the University of Canberra, and comprised skin, liver or blood snap frozen in liquid nitrogen and stored at −80°C.

Total genomic DNA was extracted by salt extraction (after Miller et al. 1988, modified by Fitz-Simmons et al. 1995). Sub-samples of skin were digested overnight at 55°C in 300 μl of buffer (40 mmol Tris, 20 mmol EDTA, 100 mmol NaCl), 20 μl of 10% SDS and 10 μl of 10 mg ml⁻¹ Proteinase K. DNA was salt extracted and precipitated with 150 μl of 7.5 mol ammonium acetate and 1 ml of 100% ethanol, washed with 70% ethanol before being re-suspended in 50 μl of water. A 4 μl subsample of each extraction was run out on a gel to confirm the success of the extraction.

PCR of the mitochondrial ND4 region (coding) used primers ND4F (5'-CAC TTA TGA TCA AAC GCT CAT GTA GAA GC-3') (Arevalo et al. 1994) and ND4intR (5'-AGG TGT TCT CGT CTT TG-3') (Alacs 2008) to yield a 640 bp DNA fragment. PCR of the mitochondrial control region (non-coding) used primers TCR500 and EMYTHR TCR500 (5'-CAC CAC CCT CCT GAA ATA ATC-3') and PS15841 (5'-CMT TGG YCT TGT AAA CC-3') (Engstrom et al. 2004) to yield a 450 bp DNA fragment. PCR products (50 μl of each sample) were precipitated with 50 μl of 20% polyethylene glycol, washed with 80% ethanol and re-suspended in 13 μl of water. The cleanup PCR products were packaged and sent to a commercial service for sequencing. Sequence data were edited using SEQUENCER V 4.1.4 (Genes Code) and aligned using CLUSTAL X (Thompson et al. 1997), with final alignment by eye.

All sequences were visually examined to eliminate the possibility that they might be nuclear mitochondrial DNA (numts). The chromatograms were clean, lacking ambiguous base signals that might have otherwise indicated presence of both mitochondrial and nuclear paralogues in a single PCR product. The coding section of the ND4 sequences for the 7 taxa were checked for the presence of premature stop codons (Coin & Durbin 2004, Hay et al. 2004, Lerat & Ochman 2005), and none was found. The amplified DNA of all 7 taxa were aligned to a known complete ND4 genome from a distantly related tortoise, *Geochelone pardalis* (GenBank accession number DQ080041; Parham et al. 2006). The nucleotides for each sequence, including for *G. pardalis*, were converted to amino acids prior to alignment. All amplified sequences were confirmed to be ND4 gene segments prior to phylogenetic analysis. While none of these approaches eliminate the possibility of nuclear
paralogues of mitochondrial sequences in our data, they at least reduce the probability.

Phylogenies for ND4 and the control region sequences were generated using maximum parsimony (MP) and maximum likelihood (ML) as implemented in PAUP V4.0b10 (Swofford 2002). For the ML analyses, the HKY85+G (Hasegawa et al. 1985) model of sequence evolution was chosen using ModelTest (Posada & Crandall 1998) prior to being implemented in Mega (Tamura et al. 2007). Trees were rooted using a nested series of outgroups. The most distantly related taxon _Chelodina rugosa_ was chosen on the basis of the previous analyses of Georges & Adams (1992, 1996). This confirmed the use of _Elseya dentata_ as a closer outgroup taxon. _Emydura macquarii_ was not found to be a suitable outgroup taxon. Robustness of the resultant trees was determined using bootstrap analyses (2000 pseudoreplicates for MP and 1000 pseudoreplicates for ML). Uncorrected p distances were calculated using Mega (Tamura et al. 2007). The specimens examined, including their collection locality and sample identification codes, are provided in Table 1.

**RESULTS**

The 640 bp ND4 sequences yielded 3 haplotypes for _Myuchelys bellii_, each differing by only a single base representing populations of the Namoi (n = 5), Gwydir (n = 4) and Border Rivers (n = 5), respectively. _M. georgesi_ from the Bellinger River (n = 5) and _M. purvisi_ from the Manning River (n = 5) each yielded a single haplotype. _M. latisternum_ from the Tweed River (n = 1) and Richmond River (n = 1) had haplo-

<table>
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<th>Stream</th>
<th>Location</th>
<th>Unique haplotypes</th>
<th>No. specimens</th>
<th>Sample identification</th>
</tr>
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<td>1</td>
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<td>UC Q19138, Q19146</td>
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<td>1</td>
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<td></td>
<td>30°41.8'S, 151°7.16'E</td>
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<td>1</td>
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<td>2</td>
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<td>1</td>
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<td>1</td>
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</table>

*Polymorphic sites
types differing by 1 bp. The 450 bp of the control region yielded 2 haplotypes for *M. bellii* differing by only a single base, and representing populations of the Namoi-Gwydir and Border Rivers, respectively. *M. georgesi* from the Bellinger River (n = 5) yielded a single haplotype, and *M. purvisi* from the Manning River (n = 5) yielded 2 haplotypes differing by only 1 bp. Again *M. latisternum* from the Tweed and Richmond Rivers had different haplotypes, differing by 3 bp. In total, 8 unique haplotypes (concatenated ND4 and control region mtDNA fragments) were identified for species of *Myuchelys*, with 11 unique haplotypes across both ingroup (*M. bellii, M. georgesi, M. latisternum* and *M. purvisi*) and outgroup taxa (*Chelodina rugosa, Elseya dentata* and *Emydura macquarii*). A matrix of uncorrected genetic p distances for all pairs of taxa, including outgroups, combining the ND4 and control region sequences, is presented in Table 1 (GenBank accession nos.: JN006511–JN006521).

The haplotype phylogeny based on the combined ND4 and control region mtDNA sequences (Fig. 2) revealed *Myuchelys bellii* as the sister taxon to *M. latisternum*, an arrangement with very strong bootstrap support from both the MP (99%) and ML (97%) analyses. *M. georgesi* and *M. purvisi*, although once a cryptic species pair, are not sister taxa; rather, *M. georgesi* is sister to the *M. bellii/latisternum* clade (MP 90%, ML 71%). Genetic distance between *M. georgesi* and *M. purvisi* was 13.5% (Table 2), as great as between any 2 of the ingroup taxa. An unexpected result was the inclusion of 1 of the 3 nested outgroup taxa, *Emydura macquarii*, among the ingroup as sister to *M. georgesi* (Fig. 2). Indeed, the genetic distances between *E. macquarii* and the species of *Myuchelys* ranged from 7.9 to 9.6%, compared to 13.0 to 13.7% for *M. purvisi* (Table 2). The currently accepted taxonomy would have *E. macquarii* as sister to *Elseya dentata* and outside the *Myuchelys* radiation (Georges & Adams 1992, 1996, Thomson & Georges 2009, Georges & Thomson 2010). For this arrangement to be consistent with our data, an additional 28 base changes are required in addition to the 546 base changes required to explain the most parsimonious topology (Swofford 2002) (Fig. 2A). In likelihood terms, the currently accepted arrangement had a log likelihood ratio of 11.2 when compared with the tree of maximum likelihood (Fig. 2B). A permutation test of the difference between the 2 topologies (1 x 10⁶ replications) indicated that this difference was highly significant (p < 0.00013). We thus interpret the finding of *E. macquarii* within the *Myuchelys* radiation as a real feature of the ND4-control region mtDNA phylogeny.

**DISCUSSION**

Multi-locus allozyme data on the delineation of Australian chelid species boundaries (sensu Georges & Adams, 1992, 1996) indicated that populations of

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**Fig. 2. Myuchelys spp. and presumptive outgroups Chelodina rugosa, Elseya dentata and Emydura macquarii.** (A) Maximum parsimony and (B) maximum likelihood trees generated using PAUP* (Swofford 2002) with concatenated sequences of ND4 (640 bp) and control region mtDNA (450 bp). Alignment gaps were treated as deletions, yielding 971 characters of which 193 were parsimony informative. Branch lengths drawn proportional to maximum likelihood estimates of genetic divergence. Drainage basin names are provided in brackets where multiple populations of the same species were sampled (see Table 1)

<table>
<thead>
<tr>
<th></th>
<th>Mybel (Border)</th>
<th>Mybel (Gwydir)</th>
<th>Mybel (Namoil)</th>
<th>Mylat (Richmond)</th>
<th>Mylat (Tweed)</th>
<th>Mygeo</th>
<th>Mypur</th>
<th>Emmac</th>
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*Myuchelys bellii* from the Namoil and Gwydir Rivers could be diagnosable as a distinct species, but they did not have samples from the Border Rivers. Since that allozyme study (George & Adams 1992), the Bald Rock Creek (Qld) populations have been identified as distinct morphologically from those of the Namoil and Gwydir Rivers (NSW), and separate species status has been proposed (Cann 1998, Wells 2002). Such distinction failed to be sustained under more rigorous morphological analysis of the 3 populations of the Endangered *M. bellii* (Fielder 2010). Additionally, the level of genetic divergence between the 3 isolated populations is equivalent to those observed between populations of *M. latisternum* from adjacent drainages (Table 1). These exceptionally low levels of mtDNA in *M. bellii* populations suggest that they should be viewed as a single taxon. Consequently, the proposition that the population in Girraween National Park (Qld) is a separate species or subspecies from those in the Gwydir and Namoil Rivers draws no support from our data. Turtles from the 3 populations occupy a single drainage basin, and although isolated to some degree, it is possible that divergence is impeded by low-frequency genetic exchange between these populations. There is at least 1 anecdotal report from the 1970s of *M. bellii* from the Macquarie Marshes (100 km downstream of its established range and below the confluence of the waters of the Namoil and Gwydir Rivers, Fig. 1), which gives credence to the proposition that such low-level exchange is ongoing. Alternatively, the extant populations of *M. bellii* might represent the outcome of recent range contraction. Recent surveys(Cann 1998, Fielder 2010, this study) looking for *M. bellii* in stream segments between known populations in the Namoil, Gwydir and the Girraween National Park populations failed to detect *M. bellii* movements. With so few data, there is no evidence whether they are isolated and do have some (minimal) gene flow, or whether they in fact have high levels of gene flow, or whether they have diverged very recently (with or without gene flow) exhibiting almost no genetic differentiation. Further investigation is required to categorically rule out these possibilities.

Consistent with earlier studies (Georges & Adams 1992, 1996), our mtDNA phylogeny establishes *Myuchelys bellii* west of the Great Dividing Range and *M. latisternum* east of the Great Dividing Range and across the north of Australia as sister taxa. There are associated morphological similarities that are interpreted as synapomorphies (Fielder 2010), including moderately to deeply serrated shell margins, dorso-ventrally depressed carapace, broad head with furrowed head shield in aged adults and pronounced neck tubercles. Also consistent with earlier studies is the separation in the phylogeny of *M. purvisi* and *M. georgesi*, which share external morphological characters (Cann 1998), yet are not sister species. Indeed a 13.5% divergence in the present mtDNA phylogeny corroborates earlier molecular studies of a 20% divergence in fixed allelic differences (Georges & Adams 1996). Their common ancestor has all 4 species of *Myuchelys* among its descendants. This is either a remarkable case of character convergence in allopatry between *M. georgesi* of the Bellinger River and *M. purvisi* of the Manning River, or these 2 species have preserved a
shared suite of plesiomorphic (retained) characters over many millions of years. This latter interpretation seems more likely, and if correct opens a unique window on the morphology of the ancestor of the sawshelled turtle radiation. This study demonstrates that genetic and morphological divergence can be uncoupled, leading to potentially conflicting interpretations of relationships among taxa if based solely on morphological data.

The observed placement of *Emydura macquarii* in the mtDNA phylogeny is a major departure from the currently accepted phylogeny and associated taxonomy (Georges & Thomson 2010). *E. macquarii* would be expected to be sister to *Elseya dentata* (McDowell 1983, Georges & Adams 1996), and outside the *Myuchelys* radiation (Georges & Adams 1992). This group has had a long and confused history. The genus *Elseya* was originally erected for *E. dentata* (the type species) and *M. latisternum* (formerly *E. latisternum*) (Gray 1867, 1872). Boulenger (1889) redefined the genus on the basis of a longitudinal shearing ridge on the maxillary triturating surface, the alveolar ridge, present only in *E. dentata*, *E. latisternum* and *E. novaeguineae* were placed in the genus *Emydura*. Furthermore, Gaffney (1977) was unable to distinguish the genera *Elseya* and *Emydura* using cranial features (though *E. latisternum* was not available to him), and later Gaffney (1979) used the 2 in synonymy. McDowell (1983) interpreted a wider range of morphological characters to indicate that the closest relative of *E. dentata* is *E. australis* (including *E. krefftii* and *E. subglobosea*) and not *E. latisternum*. He concluded that 'the generic recognition of *Elseya* seems unwarranted' and synonymised *Elseya* and *Emydura*. In contrast, Legler & Cann (1980) considered that morphological similarities between species in the 2 genera are sufficient to indicate a common ancestry, but not to warrant lumping the genera. The matter was resolved when Georges & Adams (1992) demonstrated, using allozyme electrophoresis, that the genus *Elseya* was paraphyletic, with *Emydura* coming between clades referred to today as *Elseya* and *Myuchelys* (Thomson & Georges 2009). Unlike Gaffney (1979) and McDowell (1983), Georges & Adams (1992) resolved the paraphyly by recommending a new genus for *E. latisternum* and its relatives (now *Myuchelys*). At the time, it was noted that the nearest living relative to *E. dentata* was *Emydura australis* (including *E. krefftii* and *E. subglobosea*; McDowell 1983) and not *E. latisternum*, a result supported by the allozyme data (Georges & Adams 1992, 1996).

Clearly, the outcome of the mtDNA studies further complicates the matter. In terms of the mtDNA sequences we examined, *Emydura macquarii* and *Myuchelys georgesi* are closer to each other than either is to *Elseya dentata*, and this is reflected by a sister relationship between the former 2 in the mtDNA phylogeny. If verified, this finding would require the generic names and definitions to be revisited, either by synonymising genera, erecting yet another monotypic genus, this time for *M. purvisi*, or recognising and accepting the paraphyly and retaining the existing nomenclature. We do not propose to take this action here because our data are based on mitochondrial DNA, and potentially subject to the factors that can affect any studies based on a single locus. Hybridisation between Australian chelid turtle species is widespread and there is at least 1 instance of hybridisation between *M. latisternum* and *E. macquarii* (C. Limpus pers. comm.). The possibility of historical horizontal transfer of mitochondrial genomes requires that this finding be presented as a hypothesis for testing in future with multilocus data, molecular or morphological.

Notwithstanding the taxonomic uncertainties, *Myuchelys* is a phylogenetically distinct group of chelids emerging as a priority for conservation management within the Australasian fauna. Three out of the 4 species have extremely restricted distributions, being restricted to headwaters of single drainages which makes them particularly vulnerable to ongoing threatening processes within their catchments. *M. bellii* is of particular concern. The 3 major populations of *M. bellii* are distributed across 2 jurisdictions (Qld and NSW) and primarily along stretches of river in private hands. The small Qld population (<400 individuals, Fielder 2010) from Bald Rock Creek (<10 km reach) has experienced significant habitat modification through mechanical removal of in-stream boulders and riparian vegetation and loss through impoundment of a major permanent natural waterhole. Whilst the Qld population appears healthy and stable, these threatening processes and the small population size make the question of its species status more important from a conservation point of view. A significant proportion of the restricted, but locally abundant, Namo I and Gwydir turtle populations are suffering serious eye damage and blindness from eye cataracts from unknown causes (Cann 1998). Our study has shown little evidence of divergence of the 3 populations of *M. bellii*, and as such, they should be managed as a single operational taxonomic unit. This will involve additional attention both off and on reserve to protecting the remaining refugial habitat in the headwaters of the Namo, Gwydir and Border Rivers of the Murray-Darling
catchment, as well as attention to the corridors that enable free and effective passage between these when conditions allow. The persistence and quality of wetlands like the Macquarie Marshes may prove pivotal in the long-term persistence of the 3 ‘isolated’ populations of the Endangered *M. bellii*.

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