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Deakin, Janine E., Cooper, Desmond W., Sinclair, Jennifer J., Herbert, Catherine A., Renfree, Marilyn B., and Wakefield, Matthew (2013). Towards an understanding of the genetic basis behind 1080 (sodium fluoroacetate) tolerance and an investigation of the candidate gene *ACO2*. *Australian Journal of Zoology* **61**, 69–77.

http://dx.doi.org/10.1071/ZO12108

Towards an understanding of the genetic basis behind 1080 (sodium fluoroacetate) tolerance and an investigation of the candidate gene *ACO2*

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Running title: Genetic basis of 1080 tolerance

1 Abstract

Sodium fluoroacetate, commonly referred to as 1080, is a pesticide heavily used to 2 control vertebrate pests. The development of tolerance to this poison by target species 3 is a critical concern raised by its intensive use. Tolerance to 1080 is common amongst 4 many native vertebrates in southwest Western Australia and is thought to be the result 5 of a long period of co-evolution with plant species that produce 1080 in their seeds 6 7 and flowers. Among those vertebrate species tolerant to 1080 exposure is a subspecies of the tammar wallaby (Macropus eugenii). Tammars from Western Australia are 8 9 tolerant while the subspecies present on Kangaroo Island is susceptible to 1080 exposure. The availability of genetic and genomic information, combined with a 10 distinct difference in tolerance to 1080 between subspecies, makes the tammar 11 12 wallaby an ideal species in which to study the genetic basis behind 1080 resistance. To date, research in this area has focused on a candidate gene approach. Since 1080 13 inhibits the action of the mitochondrial aconitase enzyme, the aconitase gene ACO2 14 15 was considered a prime candidate for involvement in 1080 tolerance. However, sequencing of the full-length ACO2 transcript failed to identify a sequence variant 16 between the two subspecies that would result in an amino acid change in the active 17 site of the enzyme. Future studies will need to take a genome-wide approach to 18 identify the gene(s) responsible for 1080 tolerance. 19

20

21 Introduction

Sodium fluoroacetate (FCH₂COONa), also known as 1080, is a highly toxic pesticide 22 23 intensively used for the control of introduced vertebrate pests. It is the preferred poison used in baits for pest control in Australia and New Zealand because it poses 24 less risk to non-target animals compared to other poisons such as strychnine, pindone 25 and cyanide (Fleming et al 2001). In New Zealand, 1080 has been used since the early 26 27 1950s in attempts to control the pest population of Australian brushtail possums 28 (Montague 2000) as well as the introduced Bennett's (*Macropus rufogriseus* 29 rufogriseus) and tammar wallaby (Macropus eugenii) populations (Eason et al. 2010). 1080 has been used in Australia for many years to control introduced species such as 30 rabbits (Oryctolagus cuniculus), red foxes (Vulpes vulpes) (Saunders et al. 2010), pigs 31 32 (Sus scrofa) (Twigg et al. 2005) and wild dogs (Fleming et al. 2001) and native species in Tasmania, such as common brushtail possums, Tasmanian pademelons 33 (Thylogale billardierii) and Bennett's wallabies (McIlroy 1982a). One of the concerns 34 35 raised by the intensive use of this poison is the development of resistance in target species. In fact, resistance to 1080 has already been reported for rabbits in Western 36 Australia (Twigg et al. 2002). The development of resistance in target species could 37 have major implications for vertebrate pest control programmes, as well as agriculture 38 in regions using 1080. 39

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Many native vertebrates in Western Australia are more tolerant of 1080 exposure than
their eastern Australia conspecifics. Native animals in the south-west of Western
Australia have co-existed with fluoroacetate-bearing vegetation for long periods of
time and have developed varying levels of tolerance to 1080 (Twigg *et al.* 2003).
Plants in this region belonging to the genera *Gastrolobium, Oxylobium* and *Acacia*,

produce sodium monofluoroacetate as a defence against over-browsing (Twigg and 46 King 1991). The toxic nature of these plants on livestock was recognised in the mid 47 48 1800s yet it was apparent that native vertebrate species that fed on these plant species were tolerant (Cameron 1977). Such plants are absent in eastern Australia and native 49 animals outside of Western Australia do not have 1080 tolerance. Differences in 50 51 tolerance to 1080 between populations in south-west Western Australian and other 52 parts of Australia have been reported for many species, including mammals, reptiles and birds (Oliver et al. 1977; King et al. 1978; Oliver et al. 1979; Mcilroy 1981, 53 54 1982a, b; Mead et al. 1985; Twigg and King 1989; Twigg and Mead 1990; Twigg and King 1991; Martin and Twigg 2002; Twigg et al. 2003). The genetic basis for this 55 difference in tolerance is currently unknown but it is clear that it is a genetic trait 56 (Oliver et al. 1979). 57

58

Fluoroacetate inhibits the tricarboxylic acid (TCA cycle), resulting in an accumulation 59 of citrate in plasma and tissues. Plasma citrate accumulation has been used for many 60 years to quantify the difference in fluroacetate tolerance between eastern and western 61 conspecifics and to estimate the LD_{50} (lethal dose required to kill 50% of the test 62 species) (reviewed in (Twigg and King 1991). Tolerance differences have been 63 detected between eastern and western races of reptiles, birds, as well as several 64 marsupial and eutherian mammals (Table 1). The high tolerance of native species has 65 been exploited in successful fox control programmes in Western Australia to enhance 66 conservation efforts for several threatened species, such as the Southern brown 67 bandicoot (Isoodon obesulus) (Green 2004) and the black-flanked rock wallaby 68 (Petrogale lateralis) (Kinnear et al. 1998). 69

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71 (Cooper and Herbert 2001) expressed concern over repeated use of pesticides such as 1080, making it clear that, in terms of population genetics, the potential selection for 72 resistance to a pesticide would be strong. This is an understudied area of 1080 73 74 population control programmes but, in light of the rapid detection of rodents resistant to rodenticides (Boyle 1960), and the ecological and economic consequences 1080 75 76 resistance would have for Australia and New Zealand, it is an area of research that 77 should receive more attention. Cooper also recognised the value of the tammar wallaby (Macropus eugenii) as a model species for uncovering the genetic basis 78 79 behind 1080 tolerance. Thanks in large part to his efforts to establish a pedigreed colony of tammar wallabies (McKenzie et al. 1993), this species has been extensively 80 used for research into marsupial genetics and genomics (Renfree et al. 2011). The 81 82 geographic populations of tammars, which have been isolated for at least 10,000 years, are restricted to off-shore islands and the mainland of south-west Western 83 Australia, and Kangaroo Island off the coast of South Australia. Western Australian 84 85 and Kangaroo Island populations are classified as separate subspecies. The Kangaroo Island animals are far more susceptible to 1080 than their Western Australian 86 counterparts (Oliver et al. 1979). Many genetic polymorphisms have been detected 87 between the two subspecies (Zenger et al. 2002; Wang et al. 2011a) and the genome 88 of a Kangaroo Island individual has recently been sequenced (Renfree et al. 2011). 89 90 Thus, the tammar wallaby is a good choice of species in which to investigate the 91 genetics behind 1080 tolerance.

92

93 *Elucidating the genetic basis of 1080 tolerance*

By considering the phenotype of tolerant animals we are able to deduce the likely

95 mechanisms by which tolerance is occuring. The mechanisms by which resistance to a

selective agent can occur include: reduced absorption, increased excretion, 96 detoxification, mutation of a molecular target, and metabolic pathway compensation. 97 The observations of the 1080 phenotype show that tolerance is present when 1080 is 98 administered by non-oral routes (e.g. intraperitoneal injection) to tammar wallabies, 99 western grey kangaroos (*Macropus fuliginosus*) and bush rats (*Rattus fuscipes*) 100 (Oliver et al. 1979). Species that are resistant to 1080 are also known to retain a 101 102 secondary toxicity to predators (McIlroy and Gifford 1991; Gillies and Pierce 1999). Although direct tissue distribution measurements are not available these observations 103 104 suggest that resistant species retain sufficient fluoroacetate or fluorocitrate metabolite in the consumed tissues to be toxic to the carnivour. These observations are consistent 105 with a mutation of the molecular target of the poison or a metabolic pathway 106 107 compensation rather than reduced absorption, increased excretion or detoxification being the primary mechanism of the tolerance observed in wild populations. 108 To identify the molecular basis of 1080 tolerance it is useful to understand how 109 resistance develops in a lab setting, and the broader theory of resistance evolution in 110 other contexts such as insecticide resistance. Lab based experiments have 111 demonstrated the ability to induce 1080 resistance at seven fold levels over 25 112 generations in flies (Musca domestica) and 1.8 fold over five generations in rats 113 (Tahori 1963; Howard et al. 1973). These data demonstrate that in these species 1080 114 115 resistance is a selectable trait; however, it does not indicate whether the observed resistance is monogenic or polygenic in origin. From work on insecticide resistance it 116 is suggested that acute doses that are lethal will favour single mutational events, for 117 example mutations in the active enzyme site (McKenzie and Batterham 1994). Often 118 mutations in the same gene are observed in different populations and even in different 119 species. An example of this in vertebrates would be the development of resistance to 120

anti-coagulants used to control rodents. The use of such rodenticides started in the 121 1950s and it was later that same decade that individuals resistant to the poison were 122 123 identified (Boyle 1960). This resistance has been attributed to amino acid changes in the VKORC1 gene encoding for vitamin K reductase complex subuit 1, which is a 124 target enzyme of these poisons (Rost et al. 2004). Several different mutations in this 125 gene have been detected in independent populations of rodents (Pelz et al. 2005; 126 127 Grandemange et al. 2010), all of which are capable of conferring resistance (Hodroge 128 et al. 2011). In contrast, low dose long term selection, as may occur with the natural 129 occurrence of sodium monofluoroacetate in the field, may favour a collection of more subtle changes that act together, along with the evolution of compensatory changes 130 that ameliorate the otherwise deletrious effects of resistance mutations (McKenzie & 131 Batterham, 1994). These types of mutations are often specific to particular 132 populations. 133

134

The toxicity of 1080 is the result of inhibition of aerobic production of energy of the 135 mitochondrial aconitase enzyme (EC 4.2.1.3) (Goncharov et al. 2006). This enzyme 136 catalyzes the interconversion of citrate to isocitrate in the TCA cycle. Although 137 aconitase is a mitochondrial enzyme it is encoded by a nuclear gene called aconitase 2 138 (ACO2) gene, making it a prime candidate for investigating the genetic basis of 1080 139 140 tolerance. To determine whether ACO2 plays a role in tolerance to 1080, we examined the effect of 1080 on aconitase activity in the white blood cells of tolerant 141 and susceptible subspecies of tammar wallaby as well as subspecies hybrids. White 142 blood cells were chosen because they can be obtained repeatedly without killing the 143 animals. We also sequenced the most of the ACO2 gene transcript from Western 144 Australian and South Australian tammar wallabies in order to determine if tolerance 145

- to 1080 was due to sequence differences in the candidate gene ACO2. This
- 147 investigation is a first step towards defining the genetic basis of tolerance to 1080.

149 Materials and Methods

150 *Study animals*

M.eugenii derbianus were collected from Tutanning Nature Reserve (mainland 151 Western Australia -WA), Abrolhos Island (AI) and Garden Island (GI) in the 152 southwest of Western Australia. M.eugenii decres were collected from Kangaroo 153 Island (KI) which is located in South Australia. Crosses between the two subspecies 154 were bred in captivity. The wallabies were housed in outdoor grassy yards at either 155 the Cowan Field Station, a UNSW field facility, or The University of Melbourne 156 captive colony. Water and commercial kangaroo pellets, or lucerne hay, were 157 provided ad libitum. All sampling was done with the approval of the University of 158 New South Wales Animal Care and Ethics Committee (05/25B and 05/26B) and The 159 160 University of Melbourne Animal Ethics Committee.

161

162 *Sample collection for aconitase assay*

Blood samples (8-10 mL) were collected from individuals (Total n = 36; Tutanning n = 9, Garden Island n = 3 and Kangaroo Island n = 18, and hybrids n=6) by venipuncture of a lateral tail vein using syringes containing 0.5mL Heparin (DBL[®], Melbourne, VIC, Australia) to prevent clotting. White blood cells were isolated using LeucoSep[®] separation tubes (Greiner bio-one, Frickenhausen, Germany) and Ficoll-PaqueTM PLUS (Amersham Biosciences, Uppsala, Sweden). Isolation procedures were performed according to the protocol supplied with the LeucoSep[®] separation tubes. Cell counts were conducted using a haemocytometer by viewing isolated cells
stained with a 1:2 dilution of Trypan blue (Sigma, Castle Hill, Australia) under a
microscope.

173

174 *Aconitase assay*

White cells were assayed for aconitase activity by spectrophotometrically measuring 175 the production of NADPH based on the methods of Morrison (1954) and Gardener et 176 al. (1994). After isolation, white cell concentrations were adjusted to 2-3 x 10^5 cells 177 mL⁻¹ for the aconitase assays. The cells were incubated for 1 hour at 35°C in 100mM 178 Tris buffer. The samples then underwent four rapid freeze-thaw cycles by storing 179 them for 10 minutes at -80°C and returning them to 35°C. Thawed cell solutions were 180 181 promptly assayed for aconitase activity by following the linear absorbance change at 340nm (NADPH wavelength) using a spectrophotometer (Cary 100 UV-Visible 182 spectrophotometer, Varian Inc. Australia) for 60 minutes at 22°C in a 3mL reaction 183 mix containing 36mmol Tris pH 7.4, 0.07mmol citric acid, 0.18mmol ß-nicotinamide 184 adenine dinucleotide phosphate, 1.3mmol manganese sulfate, 0.7 units isocitric 185 186 dehydrogenase (NADP+ dependent) and 7.8mM sodium fluoracetate. Positive control assays excluding sodium fluoroacetate were set up in the same manner. Samples 187 188 containing only white blood cells and sterile Milli-Q water were also assayed as a 189 negative control. The reagents in reaction mixture were all obtained from Sigma (Castle Hill, Australia). 190

191 The change in linear absorbance at 340nm can be used to calculate the amount of 192 isocitrate formed by the catalysis of isocitrate dehydrogenase. The amount of NADPH 193 produced in the reaction is stoichiometric with the amount of isocitrate produced as a

result of aconitase activity (Pallanca *et al.* 1989). One milliunit/mL⁻¹ of enzyme

195 catalyses the formation of 1 nmol/mL^{-1} of isocitrate based on the following equation:

nmol/mL⁻¹ of
$$(\Delta A_{340nm}/min \text{ Test} - \Delta A_{340nm}/min \text{ Blank})(V_F)(df)$$

isocitrate = (6.22)(V_{ISO}) x 1000

196

where V_F is the final volume of the of the assay ($V_F = 3mL$), df is the dilution factor

198 (df = 1), and V_{ISO} is the volume of isocitrate dehydrogenase used in the assay (V_{ISO} =

199 0.1mL). 6.22 is the millimolar extinction coefficient of β -NADPH at 340nm.

200

201 Sample collection for RNA extraction

Blood samples (~5mL) were collected from individuals (Tutanning n=2, Kangaroo Island n=1, Abrolhos Island n=1, KI/WA n=3, KI/GI n= 2) by venipuncture of a lateral tail vein and transferred into EDTA-containing tubes. Samples were either transferred directly into RNAlater (Ambion, Austin, TX, USA) as whole blood for transportation and storage or white blood cells were first separated from other blood components by centrifugation and then stored in RNAlater (Ambion) at -20°C until the RNA extraction process was performed.

209

210 *ACO2 transcript sequencing*

RNA was extracted from white blood cells using the RiboPure kit (Ambion)
according to the manufacturer's instructions. Any contaminating genomic DNA
present in resulting RNA was removed with by DNase treatment with the DNA-free

DNase kit (Ambion) following the manufacturer's protocol. The quality and quantity 214 of RNA extractions was determined spectrophotometically and RNA integrity was 215 tested by running all samples on a 1.2% denaturing formaldehyde agarose gel 216 (Sambrook et al. 1989). First strand synthesis of cDNA was performed on lug of total 217 RNA with GeneRacer Oligo dT primer (Invitrogen, Carlsbad, CA, USA) or random 218 hexamers using the SuperScript III Reverse Transcriptase system (Invitrogen) 219 according to the manufacturer's instructions. To ensure that there was no residual 220 contaminating genomic DNA present in the cDNA samples, a RT-negative control 221 reaction was set up for each sample where the Supercript III enzyme was excluded 222 from the first strand synthesis reaction and was used as control in all subsequent PCR 223 224 amplification experiments.

225

Rapid Amplification of cDNA Ends (RACE) was used to amplify the 5' and 3' 226 untranslated regions of the gene in one individual. Nested primers (Table 1; Figure 227 1A) were designed as close as possible to either end of the coding sequence. RACE 228 229 was performed with the GeneRacer kit (Invitrogen) and by following the 230 manufacturer's protocol. PCR amplicons were subjected to electrophoresis on a 1% agarose gel with TAE buffer and SYBR Safe DNA gene stain (Invitrogen). Resulting 231 232 bands were excised from the gel and the DNA purified using the PureLink Quick Gel Extraction kit (Invitrogen). Purified DNA was set to the Australian Genome Research 233 234 Facility (AGRF) (Brisbane, QLD, Australia) for direct sequencing.

235

236 BAC clone isolation and physical mapping

237 Overgo probes for the tammar *ACO2* gene were designed from tammar genome

sequence using the program OvergoMaker (available as a download from Washington

- 239 University Genome Sequencing Center): ACO2_A 5'-
- 240 CACTGACCTTGTCAGAGAAGATTG-3' and ACO2 B 5'-
- 241 TCCAGGTGACCATACACAATCTTC-3. These overgos were radioactively labelled
- and used to screen high density tammar wallaby BAC library filters (Me KBa;
- Arizona Genomics Institute, Tuscon, AZ, USA) using the protocol previously
- described by (Deakin et al. 2007). Resulting positive BACs were further screened
- using dot blots as previously described (Deakin et al. 2008). BAC DNA extracted
- using the Promega SV Wizard kit (Promega, Alexandria, NSW, Australia) was
- 247 labelled with SpectrumOrange (Abbott Molecular Inc., Des Plaines, IL, USA) and
- 248 hybridised to male tammar wallaby metaphase chromosomes as reported previously
- 249 (Alsop et al. 2005). Unbound probe was removed using the washing procedure
- outlined in (Deakin et al. 2008). DAPI-stained chromosome and fluorescent signal
- 251 images were captured on a SPOT RT Monochrome CCD (charge-coupled device)
- camera (Diagnostic Instruments In., Sterling Heights, MI, USA) and merged using IP
- Lab imaging software (Scanalytics Inc, Fairfax, VA, USA).
- 254
- 255 Polymorphism search within the ACO2 gene

Primers for the amplification of 5' and 3' regions and the entire ACO2 coding

sequence from all individuals were designed from sequence obtained from RACE and

- sequence available from the tammar genome project (Renfree *et al.* 2011). Primer
- sequences and product sizes are listed in Table 2 and their position within the gene for
- the six primer pairs is shown in Figure 1A. All PCR amplifications were performed in
- a 25µl reaction volume with 200ng cDNA, 0.2µM of each primer, 1X PCR Buffer
- 262 (Invitrogen), 0.2mM of each dNTP, 1.5mM MgCl₂ and 0.2µl of Platinum Taq DNA
- 263 polymerase High Fidelity (Invitrogen) and cycling conditions of an initial denaturing
 - 12

264	step of 95°C for 2min, followed by 35 cycles of 95°C for 30sec, 60°C for 30 sec,
265	72°C for 90sec, and concluded with a final extension cycle at 72°C for 10 minutes.
266	PCR products were electrophoresed on a 1% agarose gel with 1X TAE. Bands were
267	excised from the gel, DNA extracted and sent for direct sequencing at AGRF. All
268	bands were sequenced twice in both directions. Sequences were trimmed of any poor
269	quality sequence so that at least 90% of the sequence analysed had a high confidence
270	value as indicated by Sequencher software version 4.10.1 (Gene Codes Corporation,
271	Ann Arbor, MI, USA). Single nucleotidepolymorphisms (SNPs) were detected by
272	manually examining the sequence in Sequencher and were only scored as a SNP if
273	detected on both strands and/or in sequence from overlapping PCR products.
274	
275	Results and Discussion
276	Mitochrondrial aconitase tissue culture assay
276 277	<i>Mitochrondrial aconitase tissue culture assay</i> The effect of 1080 on mitochrondrial aconitase activity was examined using an <i>in</i>
276 277 278	<i>Mitochrondrial aconitase tissue culture assay</i> The effect of 1080 on mitochrondrial aconitase activity was examined using an <i>in</i> <i>vitro</i> tissue culture assay. This assay was designed to measure the production of
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With data indicative of differences in mitochondrial aconitase activity between 1080 289 tolerant and sensitive populations, the next logical step for elucidating the genetic 290 basis of tolerance was to examine the ACO2 gene for changes in amino acid sequence 291 292 in active sites of the aconitase enzyme. The ACO2 gene has been partially sequenced as part of the tammar wallaby genome project (Renfree et al. 2011). An individual 293 from Kangaroo Island was used for the sequencing project. The Ensembl genebuild 294 295 for this species (http://www.ensembl.org/Macropus eugenii/Info/Index) places ACO2 on genescaffold 8869, flanked by genes PHF5A and POLR3H as it is in most 296 297 vertebrates. However, the fragmented nature of the tammar genome sequence left gaps in the sequence, resulting in an inaccurate gene prediction. We have been able to 298 fill these sequence gaps by amplifying and sequencing the cDNA for this gene from 299 300 the tammar wallaby and provide accurate information on the structure of this gene. 301

We identified an additional 296 bp of transcript sequence not found in the tammar 302 303 genome assembly, including the first exon (36bp), 198 bp spanning exons 8 and 9 and 62bp of exon 14 (Figure 1A). The coding region of wallaby ACO2 is actually 2349bp 304 (GenBank Acession HQ646994) and encodes for a deduced 782 amino acid protein. 305 Sequence similarity at the nucleotide level with other vertebrate species ranges from 306 80% (chicken and opossum) to 85% (cow) and sequence identity at the amino acid 307 308 level ranges from 90% with chicken to 93% (opossum, human, mouse, cow). In addition, we used RACE to obtain the 5' and 3' untranslated regions of the wallaby 309 ACO2 gene. The 12bp of the 5'UTR were also not present in the tammar genome 310 sequence, most likely falling in a sequencing gap. The 3'UTR is 363bp and most of 311 this sequence is present in the genome assembly. 312

313

The location of a BAC clone (MeKBa_455G18), isolated from the tammar wallaby BAC library and confirmed to contain the *ACO2* gene, was determined using fluorescence in situ hybridisation. The BAC clone mapped to the long arm of chromosome 3 (Figure 3) in a region previously shown to contain genes from human chromosome 22 (Renfree *et al.* 2011) where the human orthologue of *ACO2* is located.

320

321 Search for polymorphisms in the ACO2 transcript

322 The entire coding region and most of the untranslated region from nine individuals

323 was sequenced. Sequence data from the individual used for the genome project

324 (Renfree *et al.* 2011) provided a second representative of a Kangaroo Island tammar

wallaby. Only seven SNPs were detected in the 2653bp of *ACO2* sequenced across all

individuals (Figure 1B and C, Table 3). All except one SNP were synonymous

327 substitutions and the only SNP to result in an amino acid change was located at

nucleotide position 2256 (either GAT – aspartic acid or AAT – asparagine). Variation

329 at this nucleotide position was not restricted to either subspecies, nor was it located

near an active site in the aconitase enzyme (Figure 1B).

331

Sequencing of *ACO2* transcripts failed to uncover any sequence variation with a likely
role in tolerance in Western Australia tammar wallabies. It could be argued that not
enough individuals were examined to completely rule out an amino acid difference
between the tammar wallaby subspecies. However, toxicology studies on these
subspecies showed that all mainland Western Australian animals survived low to high
doses of 1080 exposure (12 wallabies in total), whereas Kangaroo Island wallabies

did not survive except when given the lowest dose (Oliver *et al.* 1979). This suggests
that most of the Western Australian tammar wallaby population are resistant to 1080.

Although differences in response to 1080 appear not to be due to amino acid changes 341 342 in the ACO2 gene, changes in regulation and the quantity of aconitase produced may be responsible for tolerance. Altered regulation could be caused by variants in the 343 344 trans acting regulatory genes, or may be caused by variations in *cis* acting gene regulatory target sequences around the ACO2 gene. Regulatory sequences can be 345 346 located some distance from the gene or within the gene's introns and affect the level of transcription of the gene. Unfortunately, the depth of sequence coverage in the 347 tammar wallaby genome assembly is too low to completely cover the gene without 348 349 sequence gaps, meaning that potentially important regulatory sequences may be missing from the genome assembly. The complete sequencing of the tammar wallaby 350 BAC clone containing ACO2 could fill in any sequence gaps and facilitate the 351 352 identification of potential regulatory elements.

353

354 Future directions using genome-wide approaches

1080 tolerance may be the result of mutations in hitherto unidentified genes

interacting with ACO2, or with other genes in the TCA cycle. For these reasons, a

357 genome-wide search may be required to uncover the region(s) involved in 1080

resistance. Fortunately, a linkage map is available for the tammar wallaby which was

constructed by using crosses between the two tammar wallaby subspecies (Zenger *et*

- *al.* 2002; Wang *et al.* 2011b); This, along with other genome resources available for
- the wallaby such as the genome sequence (Renfree *et al.* 2011), physical and
- integrated maps (Deakin 2010; Wang *et al.* 2011a), will make it possible to narrow

down candidate regions or genes in future studies. In addition, the physical

localisation of *ACO2* to chromosome 3 will help determine whether *ACO2* is within a
region segregating with 1080 tolerance.

366

Linkage mapping of 1080 tolerance in crosses between tolerant and sensitive 367 populations remains one of the most promising approaches for identification of the 368 369 gene or genes involved. Hybrids between the two tammar wallaby subspecies display tolerance in 1080 in the *in vitro* mitochondrial aconitase assay. Similarly, experiments 370 371 on native rats indicate that the F1 hybrids have an intermediate phenotype (Twigg, 2003). This indicates that tolerance may be due either to a dominant or co-dominant 372 single gene, or that tolerance is a cumulative effect of multiple loci. To investigate 373 374 the inheritance pattern a substantial number of backcross animals (1080 tolerant F1's mated to sensitive population) would be required to be quantitatively phenotyped for 375 their resistance status and genotyped with microsatellites. 376

377

The tissue culture assay for 1080 resistance, while making the assumption that the resistance mechanism occurs at the cellular level, provides the advantage of being able to quantitatively determine the LC_{50} (lethal concentration) for a single animal *in vitro* while maintaining the individual in the breeding colony. This overcomes the potential issues with dose control, sensitivity and the potential effect of environmental factors in alternative approaches such as measurement of blood citrate levels in response to low doses of 1080.

385

Given the potential complexity of the inheritance, identification of the functional
variant or variants that confer 1080 resistance will remain a complex task. To identify

additional candidate genes, RNAseq analysis to compare tolerant and sensitive
populations, and to look for potential induction of genes in response to 1080, may be
a useful approach. The most promising direction remains the generation of linkage
candidate regions that will allow focused analysis of whole genome resequencing
which has become a valuable and cost effective tool since the completion of the
tammar wallaby genome.

394

395 Acknowledgements

396 Some of the research presented here represents the work Des Cooper was pursuing

towards the end of his career. His contribution to the tammar wallaby genome project,

the development of a tissue culture assay and the linkage approach represent his

399 substantial contributions to the field of 1080 tolerance.

400

We thank Kia Bailey, James Cook and Jan Nedved for help with the UNSW tammar 401 402 wallaby colony, Scott Brownlees and Bonnie Dopheide with the Melbourne University colony. We thank Becky Choi for her work on the tissue culture assay, and 403 Auda Eltahla for help with primer design and Stephen Frankenberg for in silico 404 identification of the ACO2 sequence from the wallaby genome assembly. Work on 405 1080 resistance in the tammar wallaby was supported by an Australian Research 406 407 Council Discovery Grant to DWC (DPO851844) and was also part of the work of the ARC Centre of Excellence for Kangaroo Genomics. 408

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608 Figures

Figure 1: (A) The tammar wallaby *ACO2* gene structure, including the size and
position of the 18 exons (black) and the 5' and 3' untranslated regions (light grey).
(B) Active sites (Mirel *et al.* 1998) within the enzyme are indicated by black circles
and position of SNPs detected between the ten sequenced individuals are shown. (C)
Examples of sequence chromatograms for each SNP.



tammar wallaby chromosome 3.

632Table 1: Species with detectable differences in 1080 tolerance between western

633 and eastern Australian individuals (Twigg and King, 1991).

LD_{50} given as mg 1080 kg⁻¹

Species	Common name	Western race	Eastern race
		LD ₅₀	LD ₅₀
Reptiles			
Tiliqua rugosa	Shingleback	≥ 500	214
Varanus rosenbergi	Heath monitor	200-300	40
Birds			
Phaps chalcoptera	Common bronzewing	40	25
Marsupials			
Antechinus flavipes	Yellow-footed Antechinus	12.5	3.5
Isoodon obesulus	Southern brown bandicoot	20	7
Macropus eugenii	Tammar wallaby	≥5	0.3
Trichosurus vulpecula	Common brushtail possum	125	0.75
Eutherians			
Rattus fuscipes	Bush rat	20-80	1.1
Rattus sordidus	Dusky field rat	≥5	0.3

Primer	Sequence (5' to 3')	Product	
Name		size (bp)	
5'RACE_R1	CCGAAGCCGCAGGTATGACTTGC		
5'RACE_R2	CCAGGTGACCATACACAATCTTCTCTG	241	
3'RACE_F1	TCTAACAACCTGCTGATTGG		
3'RACE_F2	CAAGAGTTTGGTCCAGTGCCA	792	
ACO2_1F	ACAAAATGGCGCCGTATAAC		
ACO2_1R	CGATCAAGTGGTCACAGTGG	402	
ACO2_2F	GCGCCGTATAACCTGCTG		
ACO2_2R	GCCCATGGTACTCGACGAT	811	
ACO2_3F	CATCTGTATCGGGGGTTGGTG		
ACO2_3R	AGGAGGTGACGATGGTGTTC	806	
ACO2_4F	GGGCTGAAATGCAAGTCTCA		
ACO2_4R	GTGGTTCCAGGGCTGCAT	817	
ACO2_5F	CACGGATCACATCTCTGCTG		
ACO2_5R	ACTGCTGCAGCTCCTTCATC	563	
ACO2_6F	ATGGGAAAGCCAATTCAGTG		
ACO2_6R	GACCTCCAGCCACCATTAAA	767	

642 Table 2: Primers used for amplification of the tammar wallaby *ACO2* gene

647 Table 3: Genotypes of Single Nucleotide Polymorphisms (SNPs) detected in the

648 ACO2 transcript sequence for each of the ten individuals tested.

- 649 KI Kangaroo Island; WA Western Australia; AI Abrolhos Islands; GI Garden
- 650 Island; KI.WA or KI.GI hybrids
- 651

	SNP Position within ACO2							
	468	546	566	810	1205	2155	2256	
KI-1	C/C	C/C	A/A	C/C	A/A	G/G	C/T	
KI-2	C/C	C/C	A/A	C/C	A/A	G/G	T/T	
WA-1	C/C	C/C	A/A	C/T	A/C	G/G	T/T	
WA-2	C/C	A/C	A/A	C/C	A/A	A/A	C/C	
AI	C/C	C/C	A/A	C/T	A/C	A/A	T/T	
KI.WA-1	A/C	C/C	A/C	C/C	A/C	G/G	T/T	
KI.WA-2	C/C	C/C	A/A	C/T	A/C	G/G	T/T	
KI.WA-3	C/C	C/C	A/A	C/C	A/C	G/G	C/T	
KI.G1-1	C/C	C/C	A/A	C/C	A/A	G/G	C/T	
KI.G1-2	C/C	C/C	A/A	C/C	A/A	G/G	C/T	