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Author for correspondence:

Lauren C. White e-mail: lauren_white@eva.mpg.de

[†]These authors contributed equally to this work.

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Global change biology

High-quality fossil dates support a synchronous, Late Holocene extinction of devils and thylacines in mainland Australia

Lauren C. White^{1,2,†}, Frédérik Saltré^{3,†}, Corey J. A. Bradshaw³ and Jeremy J. Austin¹

¹Australian Centre for Ancient DNA, School of Biological Sciences, University of Adelaide, Adelaide, South Australia 5005, Australia

²Department of Primatology, Max Planck Institute for Evolutionary Anthropology, Leipzig 04103, Saxony, Germany

³Global Ecology, College of Science and Engineering, Flinders University, GPO Box 2100, Adelaide, South Australia 5001, Australia

LCW, 0000-0001-8085-9293; FS, 0000-0002-5040-3911; CJAB, 0000-0002-5328-7741; JJA, 0000-0003-4244-2942

The last large marsupial carnivores—the Tasmanian devil (*Sarcophilis harrisii*) and thylacine (*Thylacinus cynocephalus*)—went extinct on mainland Australia during the mid-Holocene. Based on the youngest fossil dates (approx. 3500 years before present, BP), these extinctions are often considered synchronous and driven by a common cause. However, many published devil dates have recently been rejected as unreliable, shifting the youngest mainland fossil age to 25 500 years BP and challenging the synchronous-extinction hypothesis. Here we provide 24 and 20 new ages for devils and thylacines, respectively, and collate existing, reliable radiocarbon dates by quality-filtering available records. We use this new dataset to estimate an extinction time for both species by applying the Gaussian-resampled, inverse-weighted McInerney (GRIWM) method. Our new data and analysis definitively support the synchronous-extinction hypothesis, estimating that the mainland devil and thylacine extinctions occurred between 3179 and 3227 years BP.

1. Background

During the Late Pleistocene, Tasmanian devils (*Sarcophilus harrisii*) and thylacines (Tasmanian tiger or wolf, *Thylacinus cynocephalus*) were widespread over the Australian continent [1,2]. Both species subsequently became extinct on mainland Australia, only surviving into historical times on the island of Tasmania. The thylacine was hunted to extinction after European arrival [3], while devils have suffered declines of more than 80% in less than 20 years, largely due to a transmissible cancer [4]. Based on the youngest dated fossils available, both species are assumed to have become extinct on mainland Australia during the mid-Holocene (approx. 3500 years before present, BP) [5].

The cause of these extinctions is the subject of debate, with introduced dingoes, human intensification and climate change being the three main competing, but not necessarily mutually exclusive, hypotheses [3]. Debate around extinction drivers has almost always assumed that both extinctions were roughly synchronous and therefore, potentially attributable to a common cause (or set of causes). However, the reliability of many fossil ages for devils across Australia has recently been questioned [6], shifting the mainland devil's youngest reliable fossil age back to 25 500 years BP and challenging the synchronous-extinction hypothesis. The youngest fossil age of an extinct taxon is nearly always an inaccurate proxy for the final extinction date. These two dates will inevitably diverge due to incomplete sampling, taphonomic bias and uncertainty in radiometric dating [7,8]. Many statistical models have been developed to estimate extinction time (and the associated uncertainty) using the time series of fossil ages, but their accuracy varies with the mode of extinction and sampling density over time [5].

To address these issues, we combined 44 new, highquality ages for mainland devils and thylacines with existing data that met stringent quality requirements. We applied the Gaussian-resampled, inverse-weighted McInerney (GRIWM [9]) method, which incorporates both sampling density and dating errors to calculate the confidence bounds of the mainland extinction dates for both species.

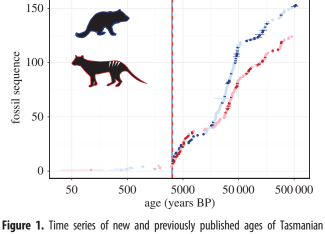
2. Material and methods

We collected less than 1 g of bone or tooth from previously undated thylacine (n = 20) and devil (n = 24) fossils from southern mainland Australia (electronic supplementary material, figure S1 and table S1) in existing museum palaeontology and mammal collections. Samples were pre-treated using ultrafiltration [10] and the collagen fraction radiocarbon-dated at the Australian National University, the University of Waikato or the Oxford Radiocarbon Accelerator Unit. We added these new ages to 129 existing mainland devil and 104 mainland thylacine museum records extracted from the FosSahul database (doi:10. 4227/05/564E6209C4FE8) [11]. We removed all unreliable ages using a set of objective criteria based on the reliability of the dating procedure used, followed by an evaluation of the confidence in the stratigraphic relationship of the dated material to the target taxon (for full details, see [6]). We calibrated all dates to calendar years (BP) using the Southern Hemisphere Calibration curve (ShCal13) from the OxCal radiocarbon calibration tool v. 4.2 (https://c14.arch.ox.ac.uk). As there is uncertainty about whether Sarcophilus laniarius was a separate, co-occurring species to Sarcophilus harrisii, or the same lineage that experienced dwarfism during the Pleistocene [12], we repeated our analyses excluding S. laniarius ages.

To estimate the mainland extinction date for these species, we followed a general guide (model-selection key in Saltré et al. [5]) for choosing the most appropriate model among the five most commonly used frequentist approaches (i.e. Solow's [7], Marshall's [13], McCarthy's [14], McInerny's [15] and GRIWM [9]) for a given series of dated fossils. Identifying the appropriateness of the models depends largely on how they treat both the probability of record occurrence and the uncertainties in record dates. GRIWM was the most appropriate method as a function of the statistical characteristics of the dataset [5] (see full analysis in electronic supplementary information and table S2). In addition to GRIWM [9], and following the same approach as Lima-Ribeiro & Diniz-Filho [16], we used the four other, wellevaluated methods regardless of the nature of sampling effort over time [17]. As these methods make different assumptions about the type of extinction (e.g. sudden versus gradual), they estimate a complete range of potential true extinction dates (see both full description and results of these approaches in the electronic supplementary material).

3. Results and discussion

The 44 new radiocarbon ages for mainland devils and thylacines (electronic supplementary material, table S1) include



devil (genus *Sarcophilus:* blue) and thylacine (genus *Thylacinus:* red) fossils from mainland Australia, showing the temporal sequence of fossils from youngest to oldest (*y*-axis) against logarithmic ages ± 1 s.d. (*x*-axis). Ages that passed quality filtering and were used to estimate extinction time are shown in dark colours, while ages that did not pass quality filtering are shown in light colours. Vertical bars represent the estimated extinction time for each species (devils: blue/solid, thylacines: red/dashed) as produced using the GRIWM approach.

the youngest, reliably dated samples for each species (devil: 3245 ± 62 years BP and thylacine: 3290 ± 49 years BP). All our new ages passed quality filtering, but only 31 of the 129 previous devil (24.0%) and 27 of 104 thylacine (26.0%) ages in the *FosSahul* database [11] met the minimum reliability criteria (A*- or A-rated only) [11]. Adding these records to the new dates produced final, high-quality datasets of 56 devil and 48 thylacine ages (figure 1). Excluding *S. laniarius* from the devil dataset left 45 reliable ages.

The continent-wide GRIWM approach produced mainland extinction estimates (T_{ext}) of 3179 years BP (95% CI: 3131–3224) for devils, and 3227 years BP (CI: 3170–3281) for thylacines (figure 1). Removing *S. laniarius* barely modified the estimate for devils ($\Delta T_{ext} = 6$ years). These results are supported by the outputs of the other statistical methods (electronic supplementary material, table S3). Our youngest reliable age for mainland devils contrasts the most recent estimate based on high-quality ages at 25 500 years BP [6]. As such, the addition of 24 new dates changes the reliable persistence timeline for this species by approximately 22 000 years.

Younger dates assessed by Rodríguez-Rey et al. [6] were mostly rejected based on inappropriate pre-treatment protocols and/or unsuitable materials used, highlighting the importance of choice in dating method. For example, all ages of bone and dentin collagen processed using ultrafiltration, ninhydrin or XAD-2 protocols or ages on individual amino acids are considered highly reliable, whereas decalcified bone or tooth dated with no information about collagen presence or those presenting collagen purification issues were discarded (see the detailed application of dating criteria for radiocarbon ages of vertebrate fossils in electronic supplementary material, table S4). Regarding ages of fossils that were not dated directly, but based on association, the strength of this association (i.e. the stratigraphic relationship between the fossil of a target species and the dated remains) was tested. Dates with no stratigraphic control or any lack of stratigraphic integrity that 2

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affects the depositional context of the target species led to rejection of that date (for more detail, see [6]).

Our results constrain the reliable dates of mainland devil and thylacine extinction to within a short (less than 50 years) period, between 3179 and 3227 years BP, which is consistent with a scenario of synchronous extinction. Synchronous extinctions have been proposed on other continents and at different time points as evidence for large-scale, common extinction drivers [18-20]. For example, analysis of the extinction chronology in North America's Pleistocene mammals suggested that a single event wiped out up to 35 genera across the continent over a 2000-year period [21]. Extending this concept to derive the most likely cause of extinctions, Cooper et al. [19] examined multiple waves of synchronous extinctions and biotic transitions across the Holarctic and found them coincident with climate warming events that likely exacerbated declines arising from human hunting. Conversely, the concurrent extinction of Australian megafauna during the Pleistocene seems to be independent of continental-scale climate change, potentially indicating a dominant human role [18,22].

Under the assumption that the mainland devil and thylacine extinctions were coincident, several studies have explored possible causes. For example, as the dingo arrived in Australia approximately 4000 years BP and never reached Tasmania, dingoes have been suspected of driving the mainland extinctions of devils and thylacines [23]. Johnson & Wroe [24] suggested that human innovation in hunting technology and more intensive use of resources could also have led to the mainland extinctions. Prowse *et al.* [3] used a modelling approach to conclude that human intensification, followed by climate change, were the most likley candidate determinants.

Other studies have avoided the assumption of synchronous extinctions by focusing on devils or thylacines separately. Letnic *et al.* [25] used morphological analyses to conclude that direct killing of thylacines by dingoes was biologically feasible and could therefore have contributed to their demise. Similarly, Brown [26] argued that climate instability associated with the onset of El Niño–Southern Oscillation (ENSO) events could have affected mainland devils, but not thylacines.

Our estimated extinction window for mainland devils and thylacines is similar to the assumed, but until now unvalidated, extinction time in most previous studies, and therefore does not challenge any aforementioned arguments, nor do our results exclude the possibility of separate or multiple causes of these extinctions. However, by supporting the assumption of synchronous extinctions with reliably dated fossil specimens, and taking into account the notion that the youngest fossil age is an inaccurate proxy for the true extinction time, our analyses provide a strong and defendable basis on which further research can build. Our understanding of these extinctions will become more complete as more palaeoclimatic, palaeoecological and archaeological data are used to uncover the biogeographic histories of these species.

Data accessibility. All new radiocarbon dates are available from the AEKOS Data Portal (doi:10.4227/05/57BA9BE3ABC2A) and have been added to the *FosSahul* database (doi:10.4227/05/564E6209C4FE8).

Authors' contributions. L.C.W. helped design the study, acquired and analysed the data and drafted the manuscript; F.S. helped design the study, analysed the data and drafted the manuscript; C.J.A.B. helped design the study, interpreted the results and helped revise the manuscript; J.J.A. helped design the study, collected the samples, interpreted the results and helped revise the manuscript. All authors gave final approval for publication.

Competing interests. We have no competing interests.

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