Scrapheap Challenge: A novel bulk-bone metabarcoding method to investigate ancient DNA in faunal assemblages

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Highly fragmented and morphologically indistinct fossil bone is common in archaeological and palaeontological deposits but unfortunately it is of little use in compiling faunal assemblages. The development of a cost-effective methodology to taxonomically identify bulk bone is therefore a key challenge. Here, an ancient DNA methodology using high-throughput sequencing is developed to survey and analyse thousands of archaeological bones from southwest Australia. Fossils were collectively ground together depending on which of fifteen stratigraphical layers they were excavated from. By generating fifteen synthetic blends of bulk bone powder, each corresponding to a chronologically distinct layer, samples could be collectively analysed in an efficient manner. A diverse range of taxa, including endemic, extirpated and hitherto unrecorded taxa, dating back to c.46,000 years BP was characterized. The method is a novel, cost-effective use for unidentifiable bone fragments and a powerful molecular tool for surveying fossils that otherwise end up on the taxonomic "scrapheap".

Fossil assemblages offer insights into past biodiversity, palaeoecology and human activities1–3. However, the accuracy of fossil identifications relies on the preservation of taxonomically significant morphological features, which are often lacking in highly fragmented remains. Over the past decade, analyses of ancient DNA (aDNA) have developed in sophistication and the breadth of contexts in which they are applied. Ancient DNA has been used to address questions of speciation, extinction and disease4–7 using a variety of substrates, including bone8, hair9 and eggshell10. However, to date, no study has attempted to use aDNA from taxonomically diverse fossils to map faunal assemblage data from a single site, largely due to the time and cost associated with generating aDNA sequences from each bone fragment.

The destructive nature of sampling also means researchers and collection managers may be reluctant to analyse valuable specimens. At the same time, most archaeological and palaeontological excavations also collect large numbers of small, morphologically indistinct bone fragments (Figure 1a). Such material is of limited use in species identifications, although it may be important for some taphonomic analyses. Taxonomically, however, it is usually destined for the analytical “scrapheap”.

It is now possible, largely due to second generation high-throughput DNA sequencing (HTS) methodologies, to genetically profile complex, heterogeneous samples (Figure 1b) in parallel, both cheaply and quickly11,12. This DNA metabarcoding13 approach to genetically unravel complex substrates via HTS, as opposed to cloning, has transformed the analysis of substrates such as sediment14,15 and faecal material16,17. To explore large HTS-generated genomic datasets from environmental samples researchers use tools that are either: 1) taxonomy-dependent, which involves searching DNA reference databases for query and reference sequence matches18, or 2) taxonomy-independent, which involves taxonomy-independent measures of sequence diversity and clustering such as Operational Taxonomic Unit (OTU) analysis or UniFrac-based methods19–22.

This study seeks to employ HTS technology to sequence and identify aDNA obtained from thousands of morphologically unidentifiable archaeological bone fragments freshly excavated from deposits at Tunnel Cave (115° 02’ E, 34° 05’ S) and Devil’s Lair (115° 04’ E, 30° 09’ S), two archaeologically and culturally significant sites
Overview of data generated. In a 2012 excavation, thousands of small bone fragments were collected by dry-sieving sediment from 15 well-dated stratigraphic units or layers at Devil’s Lair and Tunnel Cave (Figure 1c). Around 50–150 bone fragments from within each layer were each drilled for 10–15s to form 15 bulk-bone powder samples (b). Bulk-bone fragments were excavated from Devil’s Lair (DL) and Tunnel Cave (TC), two archaeologically significant sites in southwest Western Australia (c). The map used in (c) was sourced from www.openclipart.org and was modified by J.H in Adobe Illustrator.

Ancient DNA was successfully extracted from all bulk-bone powder samples, including a layer dated c.44,260–46,890 years BP (uncalibrated). The successful amplification and sequencing of DNA from all 15 layers was a rapid, cheap and effective way to assess DNA preservation at the sites (Figure 1c).

Amplicon DNA sequences (hereafter referred to as sequences) obtained from collective GS-Junior and Ion Torrent PGM sequencing runs were analysed for quality and possible chimeras. Except for ubiquitous human DNA sequences, control reactions throughout the process (described in Methods) were negative for contaminating DNA arising from laboratory processing.

Short regions within the mammalian mitochondrial 12S and 16S rRNA genes were amplified generating products of 100–104 bp and 90–96 bp respectively25. Amplification and sequencing of avian mtDNA was successful for some samples, producing either a 106–121 bp or 227–239 bp region of the avian mtDNA 12S gene25. Some cross-species reactivity was observed when using both 12S and 16S mammalian primer sets, resulting in the amplification and sequencing of avian and reptilian DNA. A targeted quantitative PCR (qPCR) and HTS approach to identify snake species was successful for a single sample.

Taxonomic identification. Mammalian 12S and 16S assays identified eight mammalian families representing 16 genera, using assignment filters chosen for this study (see Methods; Figure 2). The increase in sequencing depth afforded by the Ion Torrent PGM, as compared to the GS-Junior, did not increase the diversity of taxa identified. Mammalian taxa endemic to Australia were detected in multiple samples, in addition to taxa that have undergone significant range contraction and extirpation. The macropodid genus *Thylagale* (pademelon), provided the closest BLAST matches for many sequences across multiple samples, but to date no member of the genus has been recorded in this region. It was not possible to provide accurate taxonomic identifications for most of the Muridae sequences and for many *Macropus* sequences. While many sequences could be assigned with high confidence to a genus level, others could not be assigned beyond family or genus. A number of birds and reptiles were also identified and these have been collated at the family and genus level (Figure 2). While assignment to the species level is certainly possible in many instances a conservative approach is adopted here to showcase the approach.

Genetic biodiversity analysis. A largely taxonomy-independent approach was adopted to examine fluctuations in observed genetic diversity over time at both sites. While the taxa identified using the GS-Junior and Ion Torrent PGM were mostly congruent, coverage dependent OTU inflation, arising from homopolymer sequencing error (see Methods; Discussion) was observed. A modified OTU analysis filter was designed to reduce the influence of HTS homopolymer sequencing error26,27, by employing distance-based metrics obtained from sequence alignments, giving rise to a new method referred to here as Distance-based Taxonomic Units (DTUs).

A total of 72 DTUs were identified across all 15 samples, 23 of which were shared across multiple samples, and in some instances both archaeological sites (Figure 3). The number of DTUs fluctuates noticeably with time (Figure 4). The number of DTUs shows a notable decrease compared to the GS-Junior, did not increase the diversity of taxa identified. Mammalian taxa endemic to Australia were detected in multiple samples, in addition to taxa that have undergone significant range contraction and extirpation. The macropodid genus *Thylagale* (pademelon), provided the closest BLAST matches for many sequences across multiple samples, but to date no member of the genus has been recorded in this region. It was not possible to provide accurate taxonomic identifications for most of the Muridae sequences and for many *Macropus* sequences. While many sequences could be assigned with high confidence to a genus level, others could not be assigned beyond family or genus. A number of birds and reptiles were also identified and these have been collated at the family and genus level (Figure 2). While assignment to the species level is certainly possible in many instances a conservative approach is adopted here to showcase the approach.

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This study presents a novel HTS method using aDNA characterised from bulk-bone powder samples. It represents a powerful new approach to analyse unidentifiable fragments excavated from fossil deposits. Ancient DNA extracted from bones within a layer dated between 44,260–46,890 years BP (uncalibrated), is the oldest aDNA recovered from Australia to date. These HTS results and the initial exploration of this technique show promise for larger scale bulk-bone analyses of fossil deposits. Rapidly analysing a bulk bone sample to determine if a site is conducive to DNA preservation will be

### Figure 2 | Taxa identified in bulk-bone powder samples.

Mammals, birds and reptiles identified in each sample are listed. Samples are grouped according to site from youngest to oldest in years BP (uncalibrated), which is plotted on the same scale for both sites. The criteria used in taxonomic assignment are detailed in the Methods. Note that there is uncertainty surrounding taxonomy with regards to both Timaliidae and Cardinalidae (See Discussion). Key: √Detected using multiple primer sets; *Taxa not historically known to occur in the study region; #Sequences assigned to Rattus aligned closest to native Rattus Fuscipes (bush rat).

### Discussion

This study presents a novel HTS method using aDNA characterised from bulk-bone powder samples. It represents a powerful new approach to analyse unidentifiable fragments excavated from fossil deposits. Ancient DNA extracted from bones within a layer dated between 44,260–46,890 years BP (uncalibrated), is the oldest aDNA recovered from Australia to date. These HTS results and the initial exploration of this technique show promise for larger scale bulk-bone analyses of fossil deposits. Rapidly analysing a bulk bone sample to determine if a site is conducive to DNA preservation will be
valuable in excavations and test pits as DNA becomes increasingly incorporated into archaeological and palaeontological practices.

Even with the limited sampling, this first foray into bulk-bone analyses, has uncovered a significant amount of biological information that adds substantially to previous knowledge of the sites and surrounding biodiversity. Analysing these data in the context of DNA damage, sequencing error, incomplete reference databases and the necessary use of short DNA sequences raises numerous challenges that must be systematically addressed (see Methods) the method affords new insights into past biodiversity (Figure 2) and its temporal and spatial variation (Figures 3, 4 and 5).

Raw DNA sequences obtained from HTS platforms can be sorted and screened using a combination of filters that collectively exclude low-quality reads (Q-scores), sequences with errors in known flanking regions (adaptors, primers, and barcodes), artificial chimeric sequences and low abundance reads (see Methods). However, even sequences that pass these filters need to be interpreted with caution: the bird family Cardinalidae, which is not known to occur in Australia, is a case in point. The identification of birds also serves to illustrate the pitfalls associated with taxonomic revision. The taxonomy of the family Cardinalidae has been revised on a number of occasions, as has that of Timaliidae, which was also identified in some samples. Timaliidae has been regarded as a family consisting of Old World passerine birds, however the Australasian babblers (family: Pomatostomidae) were once within this family and the typical white-eyes (Zosterops) are disputably within this family also. The families and genera identified (Figure 2) within each of the 15 samples require further investigation to identify taxa to the species level. Nevertheless, most of the genera identified at both sites from fossil morphology were again successfully detected in

Figure 3 | DTUs shared across bulk-bone powder samples. The DTUs shared between bulk-bone powder samples, and across both Tunnel Cave (left) and Devil’s Lair (right), are shown. DTUs have been labeled with the closest BLAST family matches. Each DTU has been assigned a numeric identifier following the acronym ‘DTU’, shown in superscript. Font size is indicative of the total number of samples a DTU was detected in.

Figure 4 | Change in DTU number and composition over time at Tunnel Cave and Devil’s Lair. The fluctuation in DTU number and the change in DTU composition across samples and at both sites are plotted against the backdrop of the major climatic shift around the end of the Last Glacial Maximum (LGM). Dashed vertical line - approximate end of the LGM; Blue background – Pre-LGM; White background – LGM; Green background – Post-LGM. Median ages are plotted for each sample; dashed horizontal line indicates minimum and maximum accepted date range for each layer.
The absence of some morphologically identified taxa from the genetically-determined faunal assemblage list is most likely due to sampling bias, as the present analysis derives from deposits representing less than one percent of the volume of the original excavations. Additionally, the possibility of primer binding bias contributing to the discontinuities between both aDNA and fossil assemblage datasets cannot be excluded. In silico analysis of variation in binding sites and the use of the multiple markers attempts to identify and minimize the impact of amplification bias. Finally, inherent differences between bones in terms of the preservation and quantum of mtDNA per unit biomass may also skew results between both methods of analysis causing artifactual over-representation of some taxa relative to others. However, taxa were also identified that were not detected in any previous morphology-based analyses, particularly small mammals, birds and reptiles, all of which require highly-specialised taxonomic skills to identify, are less likely to preserve diagnostic remains, and may be poorly represented in reference collections.

A high level of confidence surrounds the bulk of the taxonomic identifications; for instance, the majority of mammalian taxa identified are locally extant or known from the fossil record. The same generally holds true for avian and reptilian taxa identifications. The detection of sequences endemic to southwest Australia, such as Tarsipes rostratus (honey possum), further supports the bona fide nature of the sequences obtained. Moreover, the detection of extirpated taxa, such as Setonix (quokka) and Sarcophilus (Tasmanian devil), as far back as c.24,000 years BP (uncalibrated) illustrates the antiquity and authenticity of the sequences, as does the detection of species whose ranges have contracted and are no longer documented at the sites, e.g. Bettongia (bettongs). There appears to be little or no environmental contamination as evidenced by the absence of any sequences from highly abundant invasive taxa including Mus musculus (house mouse) or Rattus rattus (black rat).

Whereas downward contamination may be an issue at some sites\textsuperscript{32}, Devil’s Lair contains several stratigraphical layers capped with calcite “flowstone”\textsuperscript{33} preventing the movement of fossils, and likely DNA\textsuperscript{5,23}. Whilst it is acknowledged that contamination can be cryptic and sporadic\textsuperscript{34–36}, the strict adherence to aDNA protocols\textsuperscript{37}, the use of sequence quality filters and the plausibility of the data (see Methods), greatly reduces the likelihood that contamination contributed to the data presented here.

Although most taxonomic assignments from DNA sequences confirmed previous morphological identification\textsuperscript{23}, some unexpected sequences resulted in distinct DTUs that were more difficult to assign. The issue is best exemplified by indeterminate Macropodidae sequences. It is unlikely that poor database coverage is the cause of this family-level assignment, as the Macropodidae database is nearly complete for both 16S and 12S rRNA mtDNA. In such cases sequencing error or DNA damage is also unlikely as the sequences are abundant and present across numerous samples at both sites, have passed all
quality filters, form distinct DTUs and are unlikely to be nuclear copies (Figures 2, 3 and 4). It is possible therefore that these sequences may arise from extant lineages of present-day macropodoids or indeed from extinct taxa. In some cases sequences mapped closest to species of the New Guinea forest wallaby (Dorcopsis) and the east Australian restricted pademelon (Thylagale). The presence of such ‘indeterminate’ DNA sequences in bulk-bone samples is intriguing. For example, two extinct tree-kangaroo species (genus Bothra beltrani, beltrani, have been described in caves along the Nullarbor Plain, yet tree-kangaroos of the genus Dendrolagus are only currently present in northeastern Queensland and New Guinea and were previously not thought to have occurred so far south89. It is a tantalizing prospect that ‘indeterminate’ DNA sequences could represent previously unknown species from southwest Western Australia, but it is also a problematic finding, as there is no easy way to uncover the fossils that contributed the DNA. It is likely that bulk-sampling methods such as this will generate genetically plausible taxa that lack morphological identification. Arguably a similar result has already occurred with the single Denisovan finger bone from ‘X-woman’ used to postulate a new lineage of archaic humans in Siberia80,81.

When dealing with past biodiversity and aDNA sequences from fossil assemblages, analyses that are largely independent of taxonomy will likely be crucial to mapping temporal and/or spatial variation in genetic signatures. Such an approach facilitates the use of sequences that would otherwise be labeled ‘indeterminate’, which will be commonly encountered when employing the bulk-bone HTS methodologies advocated here. While it is not possible to comprehensively analyse changes in biodiversity over time presented here from only a handful of samples such an analysis serves to illustrate how bulk-bone data could be approached. The data presented in Figures 3–5 should therefore be viewed tentatively, as further extensive replication and investigation is required to confirm any significant patternning over time.

Owing to the difficulties of definitively assigning sequences to a defined taxonomy, a modified OTU analysis (referred to as DTU), has been introduced to examine biodiversity change over time. It was clear from the initial analysis that OTU numbers were artificially inflated primarily by homopolymer error. When dealing with short sequences homopolymer errors can create a distinct OTU whereby the only difference between it and its closest OTU match is a base within a homopolymer stretch. It was observed that homopolymer-derived OTUs were more common in those samples with greater depth of sequencing coverage. To overcome this issue, an OTU alignment and Kimura 2-parameter distance matrix was adopted whereby errors in homopolymer stretches appear as gaps and homopolymer-derived OTUs collapse into a single DTU (See Methods). Whilst at these particular sites, it is a challenge to disentangle the roles of climate, DNA decay and past anthropogenic influences; shifts in DTU composition appear at the LGM and at the Holocene-Pleistocene transition (Figures 4 and 5). Furthermore, specific Macropodidae DTU analysis showed a reduction in DTU diversity and abundance over time, with a drop in diversity around the LGM (Figure 5). With these tentative patterns of biodiversity being derived from only 15 DNA extractions it is easy to conceptualize how, with adequate sampling and appropriate genetic markers, a bulk-bone sampling method will facilitate detailed mapping of faunal changes over time. Moreover, the method is cheaper than single bone approaches82–84 while augmenting traditional morphological analysis.

The bulk-bone aDNA metabarcoding method used in this study presents a new, cost effective approach to identifying bulk quantities of morphologically indistinct bone fragments that otherwise end up in the taxonomic scrapheap. From modest amounts of sieved material across multiple layers at two study sites it was possible to detect equivalent diversity as described in previous morphological analyses85. While some taxa previously identified were not detected (most noticeably Macropus species), the converse was also true. This method is by no means an attempt to supplant traditional morphological approaches to taxonomic identification and analysis. Rather, it complements these approaches and by means of DTU analysis indicates changes in genetic diversity through time. Besides improving the identification of fossil assemblages the method allows researchers to rapidly assess the DNA preservation potential of freshly excavated material, which will vary from site to site. The approach will be equally applicable to archaeological and palaeontological sites, providing snapshots of past faunal diversity and human subsistence in both taxonomic dependent and independent ways. As such, it is anticipated that a bulk-bone approach will become a valuable part of the archaeological and palaeontological toolkit.

Methods
Sample collection and processing. Thousands of indistinct bone fragments were collected from both Tunnel Cave and Devil’s Lair during excavations in February 2012. Approximately 150 L (0.15 m³) of sediment was analysed at both sites. Sediment was dry-sieved on site, using 2 mm and 5 mm sieves, and bagged according to well-defined and dated stratigraphical layers86. Each bagged sample was screened for bone fragments off-site, which were kept in groupings according to the layers in which they were found. Fifteen bulk-bone samples representing fifteen layers were screened; eight from Tunnel Cave, covering a period from 24,300 years BP (uncalibrated), and seven from Devil’s Lair, covering a period from 6,200–46,890 years BP (uncalibrated)87. Small sections of the bones within each layer (typically 50–150 bones) were drilled (Dremel 114 drill bits) for a few seconds each and approximately equal amounts of drilled material from each bone fragment within a single layer was combined to form a ‘bulk-bone powder’. Owing to inherent differences in the amount of DNA per unit of biomass between species and differential DNA preservation between individual bones, over-representation of certain bone material in terms of DNA amplexon sequences is unavoidable.

DNA extraction and screening. All laboratory work was conducted in keeping with standard aDNA protocols88. Approximately 1 g of bulk-bone powder from each sample, including a blank extraction control, was digested overnight on a lab rotator at 55 °C in 5 mL of digestion buffer containing: 2.5 mL EDTA (0.5 M), 0.1 mL Tris-HCl (1 M), 5 mg Proteinase K powder, 50 μL DTT (1 M), 50 μL Sds and made up to a final volume of 5 mL using EDTA. DNA digests were centrifuged at 6,000 rpm for 10 mins and supernatant was transferred to a clean 2 mL eppendorf tube and PBi buffer (Qiagen) for 50 μL using AMICON 30,000 MWCO columns (Millipore) as per the manufacturer’s instructions. Each concentrate was transferred to a clean 2 mL eppendorf tube and Pbi buffer (Qiagen) totalling 250 μL (i.e. 5% the volume of concentrate) was added. Each 300 μL Pbi/concentrate mix was subsequently transferred to Qiagen silica spin columns and centrifuged at 13,000 rpm. Columns were washed with 700 μL of AW1 followed by AW2. A final dry spin at 13,000 rpm for 1 min followed. DNA was eluted from the columns in 60 μL EB with a 1 min incubation at room temperature prior to centrifugation at 13,000 rpm for 1 min.

Extracts were screened for amplifiable mtDNA using multiple primer sets via qPCR with three concentrations – undiluted, 1/10 and 1/50. Extracts were screened for mammalian mtDNA using 125A/O and 165mmn mammalian primers, set designed to amplify a small region within mammalian 125 and 165 mitochondrial genes respectively89,90. Extracts were also tested for avian mtDNA using 125A/E and 125A/H primer sets, designed to amplify a short and slightly longer overlapping region of the avian mitochondrial 12S gene respectively91. Finally, extracts were tested for snake mtDNA using the following primers: 12s_irnRNA_F1_S AAAGTATAGCACTGAAAATGC and 12s_R1_Snake GTTGGCCTGTATACCGCCTCCG, designed to amplify a short region within the 125 mitochondrial gene. Each qPCR reaction was made up to a total volume of 25 μL, containing 1× PCR Gold Buffer (Applied Biosystems), 2.5 mM MgCl₂ (Applied Biosystems), 0.4 mg/mL BSA (Fisher Biotech, Aus), 0.25 mM of each dNTP (Astral Scientific, Aus), 0.4 μM forward primer, 0.4 μM reverse primer (0.25 μM AmpliTag Gold (Applied Biosystems), 0.6 μM SYBR Green (1:2,000, Life Sciences gel stain solution) and 2 μL DNA extract. Quantitative PCR cycling conditions for the 125A/O and snake 125 qPCR assays were as follows: initial heat denaturation at 95 °C for 5 mins, followed by 50 cycles of 95 °C for 30 s, 55 °C for 30 s (annealing step), 72 °C for 45 s followed by a 1 °C melt curve and final extension at 72 °C for 10 mins. Cycling conditions for 165mmn, 125A/E and 125A/H assays were the same as for the 125A/O assay, except the annealing temperature, which was 57 °C in each case. For each qPCR assay, DNA extraction, negative PCR reagent and positive DNA template controls were included.

DNA sequencing. DNA extracts that successfully yielded DNA of sufficient quality, free of inhibition, as determined by initial qPCR screening92, were prepared for amplicon sequencing. DNA extracts successful for all primer sets were sequenced on GS Junior. Additional, separate, amplicon sequences were generated for extracts using mammalian 125A/O and 165mmn primer sets for sequencing on Life Technologies’ Ion Torrent Personal Genome Machine (PGM).

For each primer set, DNA extracts were assigned a unique DNA tag93. Each sample was tagged at both the 5′ and 3′ end of the target sequence using separate tags at both
ends, resulting in a unique forward and reverse tag combination for each sequence. Independent tagged qPCRs for all samples, across all primer sets, were carried out in 25 μL reactions with reaction components and cycling conditions as described in 'Methods: DNA extraction and screening'. Tagged qPCR amplicons were generated in triplicate and combined, thus minimizing the effects of PCR stochasticity on low-template samples, purified using Agencourt AMPure XP PCR Purification Kit (Beckman Coulter Genomics, NSW, Aus), as per manufacturer’s instructions and eluted in 40 μL H₂O. Purified amplicons were pooled to form separate sequencing libraries according to primer set used and sequencing platform. GS-Junior libraries were quantified using qPCR to determine an appropriate volume of library for sequencing (described in Murray et al. 2011). Each 25 μL reaction contained 12.5 μL ABI Power SYBR master mix (Applied Biosystems), 0.4 μM A-adaptor primer, 0.4 μM B-adaptor primer, 8.5 μL H₂O and 2 μL pooled library, with the following cycling conditions: 95°C for 5 mins; 40 cycles of 95°C for 15 s, 60°C for 1 min followed by a 1°C melt curve. The appropriate library volume for use on the Ion Torrent PGM was determined using a Bioanalyzer 2100 (Agilent). For each tagged qPCR, negative controls were included and each tagged qPCR for which DNA had been found was quantified using the Bioanalyzer 2100 (Agilent) to determine an appropriate volume of library for sequencing. The GS-Junior libraries were then pooled to form the final pool for each sample, every sequence assigned to the remaining clusters were queried against sequences or total number of sequences within the most abundant cluster) was made error. Additionally, the decision to class clusters as being in low abundance with cluster selected as the representative sequence. To reduce noise associated with sequencing error, low abundant clusters, classed as those that occur at less than 1% of the total number of unique sequences when clustered at 100% sequence identity, were removed from the dataset. While the selection of a 1% cut-off is somewhat arbitrary, it should negate the possibility of clusters remaining that are the result of sequencing error. Additionally, the decision to class clusters as being in low abundance with respect to the total number of unique sequences (as opposed to total number of sequences or total number of sequences within the most abundant cluster) was made to minimize the effects of preferential DNA preservation and/or amplification. For each sample, every sequence assigned to the remaining clusters and were queried against the NCBI GenBank nucleotide database using BLASTn47 in YABIP, enabling taxonomic identification. Sequences were searched without a low complexity filter, with a gap penalties existence of five and extension of two, expected alignment value less than 1e-10 and a word count of seven. The BLASTn results obtained were imported into MEGA5 52 where a distance matrix between OTUs alignments were imported into MEGA5 52 where a distance matrix between OTUs was constructed. The species similarity between samples was constructed using the unweighted pair group method cluster analysis (UPGMA) method47 in USEARCH and were compared with the NCBI GenBank nucleotide database using BLASTn 48 to identify sequences assigned to a genus. Owing to the difficulties in assigning taxa beyond the genus level, sequences assigned to the NCBI GenBank nucleotide database were discarded from taxonomic analysis. Sequences with percentage similarity to a query sequence, such sequences were moved higher up the taxonomic rankings. When the validity of filters and hard percentage cut-offs are always debatable, those chosen in the analysis of this dataset seemed to afford the best balance when accounting for low template amounts and post-mortem damage on short adnA fragments.

Genetic biodiversity analysis. Cognisant of the difficulties associated with assigning sequences to lower taxonomic levels, a modified form of OTU analysis was applied to the 16S rRNA gene sequences obtained in this study. This allowed changes in observed genetic diversity over time at both sites to be investigated independently of the above taxonomic classification. Sequences within each sample were clustered at 97% identity, filtered and representative sequences were selected as detailed in Methods: Sequence Identification. Representative sequences within each sample were aligned in Geneious using MAFFT’s G-INS-I algorithm and default parameters5. MAFFT alignment were imported in MEGA5 where a distance matrix between OTUs were calculated using a Kimura 2-parameter model5, with the sequence containing gaps and missing data ignored. OTUs less than 3% divergent from each other were collapsed into a single OTU. This serves the purpose of reducing the influence of HTS homopolymer sequencing error5 by collapsing multiple homopolymer-derived OTUs into a single OTU, as errors in homopolymer stretches appear as gaps and are not included in the calculation of the distance matrix. Whilst this is first and foremost a largely taxonomic-independent analysis it is still nonetheless useful to identify coarsely to which family each OTU belongs, as this gives an idea of the diversity of OTUs within specific families. As such, all OTUs were searched against the NCBI GenBank nucleotide database using BLASTn5 to identify the family to which this OTU could be easily assigned. For the faunal specific Macropodidae OTU analysis the same method as above was followed except that only sequences assigned to Macropodidae were selected.

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**Author contributions**

D.C.M., M.B. and J.H. designed the experiments. D.C.M., J.H., N.W., D.H. and J.D. excavated and prepared samples. D.C.M., J.H., M.I.B., D.H. and R.A. contributed to HTS data generation and bioinformatics. J.D. provided stratigraphic interpretations and J.H. provided fossil and taxon interpretations. D.C.M. and M.B. wrote the paper with assistance from all co-authors.

**Additional information**

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