

DATA NOTE

The genome sequence of the European water vole, *Arvicola*amphibius Linnaeus 1758 [version 1; peer review: 3 approved]

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Abstract

We present a genome assembly from an individual male *Arvicola amphibius* (the European water vole; Chordata; Mammalia; Rodentia; Cricetidae). The genome sequence is 2.30 gigabases in span. The majority of the assembly is scaffolded into 18 chromosomal pseudomolecules, including the X sex chromosome. Gene annotation of this assembly on Ensembl has identified 21,394 protein coding genes.

Keywords

Arvicola amphibius, European water vole, genome sequence, chromosomal



This article is included in the Tree of Life gateway.

Open Peer Review Approval Status 🗡 🏏 🗸 version 1 24 Jun 2021 1. **Petr Kotlik** . Institute of Animal Physiology and Genetics of the Czech Academy of Sciences, Liběchov, Czech Republic 2. Alfonso Balmori-de la Puente , Institute of Evolutionary Biology (CSIC-UPF), Barcelona, Spain **Lídia Escoda** , Institute of Evolutionary Biology (CSIC-UPF), Barcelona, Spain 3. **Fahad Alqahtani** , King Abdulaziz City for Science and Technology, Riyadh, Saudi Arabia Any reports and responses or comments on the

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article can be found at the end of the article.

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Competing interests: |. Threlfall was an employee of F1000Research up until |anuary 2021.

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Species taxonomy

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Myomorpha; Muroidea; Cricetidae; Arvicolinae; Arvicola; Arvicola amphibius Linnaeus 1758 (NCBI:txid1047088).

Introduction

The European water vole, Arvicola amphibius Linnaeus 1758, is a small semi-aquatic mammal that lives on the banks of freshwater water courses and in wetlands. A. amphibius is native to Europe, west Asia, Russia and Kazakhstan. While the IUCN Red List of Threatened Species reports that A. amphibius is of "least concern" worldwide, populations in the United Kingdom have declined to such an extent that the species is considered nationally endangered (Mathews & Harrower, 2020) owing to habitat loss and predation by the American mink, Neovison vison, an invasive alien species. An estimate by Natural England put the 2018 UK population of A. amphibius at 132,000, down from 7.3 million in 1990 (Strachan, 2004). Water voles are absent from Ireland. There have been a number of conservation projects in the UK aimed at supporting populations of A. amphibius, including efforts at habitat restoration and to control the population of American mink (Bryce et al., 2011). There are also efforts to reintroduce the water vole in a number of restored urban and wild habitats. This genome sequence will be of use as a reference for researchers that wish to assess the population genomics of A. amphibius and manage reintroductions.

Genome sequence report

The genome was sequenced from a single male A. amphibius collected from the Wildwood Trust, Herne Common, Kent, UK. A total of 45-fold coverage in Pacific Biosciences singlemolecule long reads (N50 20 kb) and 52-fold coverage in 10X Genomics read clouds (from molecules with an estimated N50 of 155 kb) were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. The final assembly has a total length of 2.298 Gb in 216 sequence scaffolds with a scaffold N50 of 138.7 Mb (Table 1). The majority, 99.4%, of the assembly sequence was assigned to 19 chromosomal-level scaffolds, representing 17 autosomes (numbered by sequence length apart from chromosome 12, which is larger because the previous version of the assembly, mArvAmp1.1, mistakenly labelled this as two separate chromosomes), and the X sex chromosome (Figure 1-Figure 4; Table 2). The assembly has a BUSCO (Simao et al., 2015) v5.0.0 completeness of 96.1% using the mammalia_odb10 reference set. While not fully phased, the assembly deposited is of one haplotype. Contigs corresponding to the second haplotype have also been deposited.

Gene annotation

The Ensembl gene annotation system (Aken *et al.*, 2016) was used to generate annotation for an earlier version of the *Arvicola amphibius* assembly (GCA_903992535.1). Annotation was created primarily through alignment of transcriptomic data to the genome, with gap filling via protein to-genome

Table 1. Genome data for *Arvicola amphibius*, mArvAmp1.2.

Project accession data		
Assembly identifier	mArvAmp1.2	
Species	Arvicola amphibius	
Specimen	mArvAmp1	
NCBI taxonomy ID	txid1047088	
BioProject	PRJEB39550	
BioSample ID	SAMEA994740	
Isolate information Male; blood sample		
Raw data accessions		
PacificBiosciences SEQUEL I	ERX3146757-ERX3146763	
10X Genomics Illumina	ERX3163119-ERX3163121, ERX3341539-ERX3341546	
Hi-C Illumina	ERX3338011, ERX3338012	
BioNano	ERZ1392829	
Genome assembly		
Assembly accession	GCA_903992535.2	
Accession of alternate haplotype	GCA_903992525.1	
Span (Mb)	2,298	
Number of contigs	1,085	

Contig N50 length (Mb)	5.4
Number of scaffolds	216
Scaffold N50 length (Mb)	138.7
Longest scaffold (Mb)	199.8
BUSCO* genome score	C:96.1%[S:94.1%,D:2.0%],F:0.8%,M:3.1%,n:9226
Genome annotation	
Number of protein-coding genes	21,394
Average length of protein-coding gene (bp)	1,700
Average number of exons per gene	11
Average exon size (bp)	208
Average intron size (bp)	4,995

^{*} BUSCO scores based on the mammalia_odb10 BUSCO set using v5.0.0. C= complete [S= single copy, D=duplicated], F=fragmented, M=missing, n=number of orthologues in comparison. A full set of BUSCO scores is available at https://blobtoolkit.genomehubs.org/view/Arvicola%20amphibius/dataset/CAJEUG02/busco.

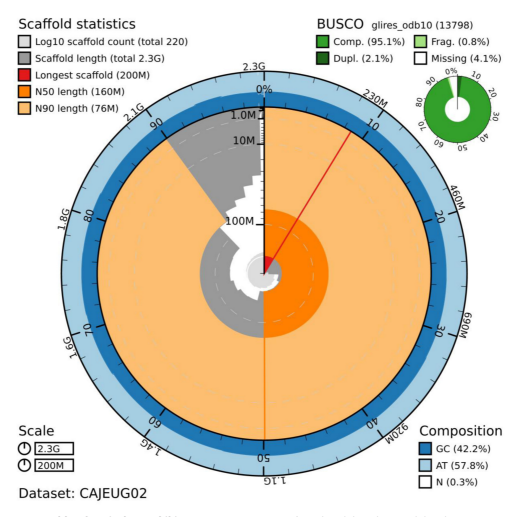


Figure 1. Genome assembly of *Arvicola amphibius***, mArvAmp1.2: metrics.** The BlobToolKit Snailplot shows N50 metrics and BUSCO gene completeness. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/Arvicola%20amphibius/dataset/CAJEUG02/snail.

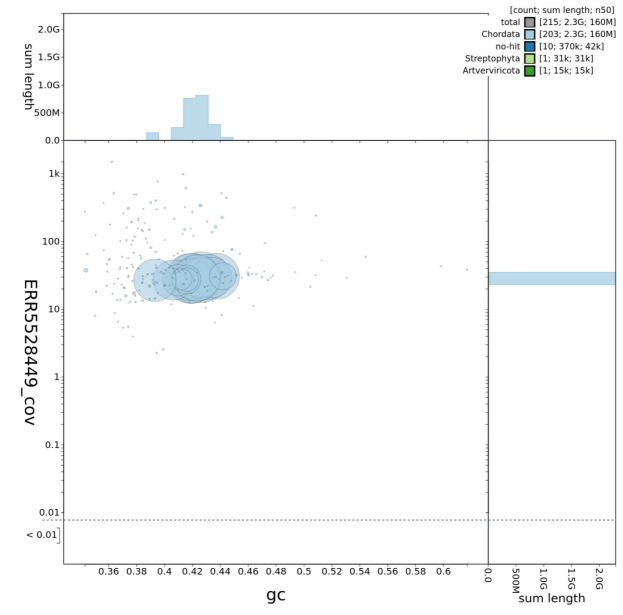


Figure 2. Genome assembly of *Arvicola amphibius***, mArvAmp1.2: GC coverage.** BlobToolKit GC-coverage plot. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/Arvicola%20amphibius/dataset/CAJEUG02/blob.

alignments of a select set of vertebrate proteins from UniProt (UniProt Consortium, 2019) and coordinate mapping of GENCODE (Frankish *et al.*, 2019) mouse reference annotations via a pairwise whole genome alignment. The resulting Ensembl annotation includes 34,750 transcripts assigned to 21,394 coding and 2,252 non-coding genes (Arvicola amphibius - Ensembl Rapid Release).

Methods

A blood sample was taken from a live male A. amphibius specimen that was part of the captive breeding population of Wildwood Trust, Herne Common, Kent, UK (latitude 51.33181, longitude 1.11443). DNA was extracted using an agarose plug

extraction from a blood sample following the Bionano Prep Animal Tissue DNA Isolation Soft Tissue Protocol. Pacific Biosciences CLR long read and 10X Genomics read cloud sequencing libraries were constructed according to the manufacturers' instructions. Sequencing was performed by the Scientific Operations DNA Pipelines at the Wellcome Sanger Institute on Pacific Biosciences SEQUEL I and Illumina HiSeq X instruments. Hi-C data were generated using the Dovetail v1.0 kit and sequenced on HiSeq X. Ultra-high molecular weight DNA was extracted using the Bionano Prep Animal Tissue DNA Isolation Soft Tissue Protocol and assessed by pulsed field gel and Qubit 2 fluorimetry. DNA was labeled for Bionano Genomics optical mapping following the

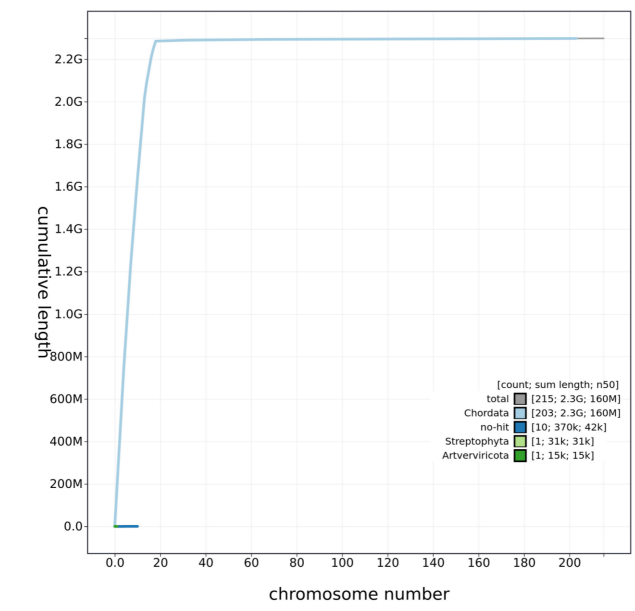


Figure 3. Genome assembly of *Arvicola amphibius***, mArvAmp1.2: cumulative sequence.** BlobToolKit cumulative sequence plot. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/Arvicola%20amphibius/dataset/CAJEUG02/cumulative

Bionano Prep Direct Label and Stain (DLS) Protocol and run on one Saphyr instrument chip flowcell.

Assembly was carried out following the Vertebrate Genome Project pipeline v1.6 (Rhie et al., 2020) with Falcon-unzip (Chin et al., 2016), haplotypic duplication was identified and removed with purge_dups (Guan et al., 2020) and a first round of scaffolding carried out with 10X Genomics read clouds using scaff10x. Hybrid scaffolding was performed using the BioNano DLE-1 data and BioNano Solve. Scaffolding with Hi-C data (Rao et al., 2014) was carried out with SALSA2 (Ghurye et al., 2019). The Hi-C scaffolded assembly was polished with arrow using the PacBio data, then polished with the 10X Genomics

Illumina data by aligning to the assembly with longranger align, calling variants with freebayes (Garrison & Marth, 2012) and applying homozygous non-reference edits using beftools consensus. Two rounds of the Illumina polishing were applied. The assembly was checked for contamination and corrected using the gEVAL system (Chow et al., 2016) as described previously (Howe et al., 2021). Manual curation was performed using evidence from Bionano (using the Bionano Access viewer), using HiGlass and Pretext, and by taking marker data and inspecting 10X barcode overlap using longranger. Figure 1–Figure 3 were generated using BlobToolKit (Challis et al., 2020). Table 3 contains a list of all software tool versions used, where appropriate.

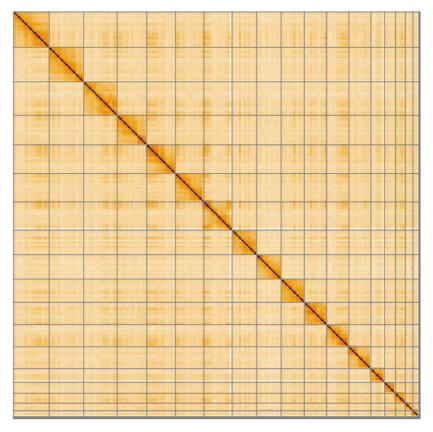


Figure 4. Genome assembly of *Arvicola amphibius***, mArvAmp1.2: Hi-C contact map.** Hi-C contact map of the mArvAmp1 assembly, visualised in HiGlass.

Table 2. Chromosomal pseudomolecules in the genome assembly of *Arvicola amphibius*, mArvAmp1.2.

INSDC accession	Chromosome	Size (Mb)	GC%
LR862380.1	1	200.53	42.7
LR862381.2	2	193.96	41.9
LR862382.1	3	189.60	42.0
LR862383.1	4	161.33	43.8
LR862384.1	5	160.72	42.6
LR862385.2	6	158.92	43.1
LR862386.1	7	138.66	41.9
LR862388.1	8	131.41	42.0

INSDC accession	Chromosome	Size (Mb)	GC%
LR862389.2	9	125.83	43.4
LR862390.1	10	125.09	42.6
LR862391.2	11	123.99	40.7
LR862392.2	12	166.75	42.3
LR862393.1	13	75.71	41.0
LR862394.1	14	63.16	41.6
LR862395.1	15	55.45	44.2
LR862397.2	17	42.65	41.6
LR862398.1	18	33.21	41.2
LR862387.1	X	137.70	39.3

Table 3. Software tools used.

Software tool	Version	Source
Falcon-unzip	falcon-kit 1.8.0	(Chin et al., 2016)
purge_dups	1.2.3-b542dbf	(Guan et al., 2020)
SALSA2	2.2-14-g974589f	(Ghurye et al., 2019)
scaff10x	4.2	https://github.com/wtsi-hpag/Scaff10X
Bionano Solve	3.3_10252018	N/A
arrow	gcpp 1.9.0-SL-release- 8.0.0+1-37-gd7b188d	https://github.com/PacificBiosciences/GenomicConsensus
longranger align	2.2.2	https://support.10xgenomics.com/genome-exome/software/pipelines/latest/advanced/other-pipelines
freebayes	1.3.1-17-gaa2ace8	(Garrison & Marth, 2012)
bcftools consensus	1.9-78-gb7e4ba9	http://samtools.github.io/bcftools/bcftools.html
gEVAL	N/A	(Chow et al., 2016)
HiGlass	1.11.6	(Kerpedjiev et al., 2018)
PretextView	0.0.4	https://github.com/wtsi-hpag/PretextMap
BlobToolKit	2.5	(Challis et al., 2020)

Data availability

Underlying data

European Nucleotide Archive: Arvicola amphibius (European water vole) genome assembly, mArvAmp1. Accession number PRJEB39550.

The genome sequence is released openly for reuse. The *Arvicola amphibius* genome sequencing initiative is part of the Wellcome Sanger Institute's "25 genomes for 25 years" project. It is also part of the Vertebrate Genome Project (VGP) ordinal references

programme and the Darwin Tree of Life (DToL) project. All raw data and the assembly have been deposited in the ENA. The genome will be annotated and presented through the Ensembl pipeline at the European Bioinformatics Institute. Raw data and assembly accession identifiers are reported in Table 1.

Acknowledgements

We thank Mike Stratton and Julia Wilson for their continuing support for the 25 genomes for 25 years project.

References

Aken BL, Ayling S, Barrell D, et al.: **The Ensembl Gene Annotation System.** Database (Oxford). 2016; **2016**: baw093.

PubMed Abstract | Publisher Full Text | Free Full Text

Bryce R, Oliver MK, Davies L, et al.: Turning Back the Tide of American Mink Invasion at an Unprecedented Scale through Community Participation and Adaptive Management. Biological Conservation. 2011; 144(1): 575–83. Publisher Full Text

Challis R, Richards E, Rajan J, et al.: BlobToolKit - Interactive Quality
Assessment of Genome Assemblies. G3 (Bethesda). 2020; 10(4): 1361–74.
PubMed Abstract | Publisher Full Text | Free Full Text

Chin CS, Peluso P, Sedlazeck FJ, et al.: Phased Diploid Genome Assembly with Single-Molecule Real-Time Sequencing. Nat Methods. 2016; 13(12): 1050–54. PubMed Abstract | Publisher Full Text | Free Full Text

Chow W, Brugger K, Caccamo M, et al.: gEVAL — a Web-Based Browser for Evaluating Genome Assemblies. Bioinformatics. 2016; 32(16): 2508–10. PubMed Abstract | Publisher Full Text | Free Full Text

Frankish A, Diekhans M, Ferreira AM, et al.: GENCODE Reference Annotation

for the Human and Mouse Genomes. Nucleic Acids Res. 2019; 47(D1): D766–73. PubMed Abstract | Publisher Full Text | Free Full Text

Garrison E, Marth G: Haplotype-Based Variant Detection from Short-Read Sequencing. arXiv:1207.3907. 2012.

Ghurye J, Rhie A, Walenz BP, et al.: Integrating Hi-C Links with Assembly Graphs for Chromosome-Scale Assembly. PLoS Comput Biol. 2019; 15(8):

PubMed Abstract | Publisher Full Text | Free Full Text

Guan D, McCarthy SA, Wood J, et al.: Identifying and Removing Haplotypic Duplication in Primary Genome Assemblies. Bioinformatics. 2020; 36(9): 2016.

PubMed Abstract | Publisher Full Text | Free Full Text

Howe K, Chow W, Collins J, et al.: Significantly Improving the Quality of Genome Assemblies through Curation. Gigascience. 2021; 10(1): giaa153. PubMed Abstract | Publisher Full Text | Free Full Text

Kerpedjiev P, Abdennur N, Lekschas F, et al.: HiGlass: Web-Based Visual

Exploration and Analysis of Genome Interaction Maps. *Genome Biol.* 2018; 19(1): 125.

PubMed Abstract | Publisher Full Text | Free Full Text

Mathews F, Harrower C: **IUCN - compliant Red List for Britain's Terrestrial Mammals.** Assessment by the Mammal Society under contract to Natural England, Natural Resources Wales and Scottish Natural Heritage. Natural England, Peterborough, 2020.

Reference Source

Rao SSP, Huntley MH, Durand NC, et al.: A 3D Map of the Human Genome at Kilobase Resolution Reveals Principles of Chromatin Looping. Cell. 2014; **159**(7): 1665–80.

PubMed Abstract | Publisher Full Text | Free Full Text

Rhie A, McCarthy SA, Fedrigo O, et al.: Towards Complete and Error-Free

Genome Assemblies of All Vertebrate Species. *bioRxiv.* 2020; 2020.05.22.110833.

Publisher Full Text

Simão FA, Waterhouse RM, Ioannidis P, et al.: BUSCO: Assessing Genome Assembly and Annotation Completeness with Single-Copy Orthologs. *Bioinformatics*. 2015; **31**(19): 3210–12.

PubMed Abstract | Publisher Full Text

Strachan R: Conserving Water Voles: Britain's Fastest Declining Mammal. Water Environ J. 2004; 18(1): 1–4.

Publisher Full Text

UniProt Consortium: **UniProt: A Worldwide Hub of Protein Knowledge.** *Nucleic Acids Res.* 2019; **47**(D1): D506–15. **PubMed Abstract** | **Publisher Full Text** | **Free Full Text**

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Version 1

Reviewer Report 23 May 2022

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Fahad Alqahtani 🗓



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This is an important study, and the authors have sequenced, assembled, and annotated a single male Arvicola amphibious (European water vole). According to the Wildlife Trusts, this species is endangered in Great Britain and on the England Red List for Mammals. From 1990 to 2018, the UK population of the European water vole dropped from 7.3 million to 132 hundred thousand. This population loses about 98% of its species in less than 30 years. Next-generation sequencing technologies are an efficient approach to generating life history and demographic data with respect to the management of endangered wildlife. Here, the authors have used different sequencing technologies (PacificBiosciences SEQUEL I, 10X Genomics Illumina, Hi-C Illumina, and BioNano) and generated a high-quality chromosomal-level assembly of this important species.

The manuscript is well written and well designed, and the results are clearly presented. The article adds much to the scientific community.

Is the rationale for creating the dataset(s) clearly described?

Yes

Are the protocols appropriate and is the work technically sound?

Are sufficient details of methods and materials provided to allow replication by others?

Are the datasets clearly presented in a useable and accessible format?

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Applied algorithms in the bioinformatics field. Mitochondrial assembly and Mitochondrial haplogroup assignment.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 17 May 2022

https://doi.org/10.21956/wellcomeopenres.18475.r50221

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Alfonso Balmori-de la Puente 🗓

Institute of Evolutionary Biology (CSIC-UPF), Barcelona, Spain

Lídia Escoda 🗓

Institute of Evolutionary Biology (CSIC-UPF), Barcelona, Spain

This manuscript presents the annotated genome sequence of the European water vole (*Arvicola amphibius*), a small semiaquatic rodent distributed across Europe and Asia. Water vole populations of the genus *Arvicola* have a complex evolution with fossorial and semi-aquatic ecological types (ecotypes), thus this genome sequence can be very convenient to study ecological adaptations in rodents.

The report is well structured and clearly defined. However, there are some parts in the introduction that need to be better clarified.

The controversial taxonomic status of this genus, specifically between *A. amphibius* and its sister species *A. scherman*, and the complex genetic structure found in Great Britain is not properly assessed in the introduction. In addition, water voles of the genus *Arvicola* have broad ecological variability that should be better explained. Based on this, the species and the ecotype analyzed should be identified. In addition, the postglacial colonization events of water voles in the United Kingdom might be explained in more detail. All of these aspects will facilitate the future applications of the specimen sequenced for the conservation of the species.

Is the rationale for creating the dataset(s) clearly described?

Yes

Are the protocols appropriate and is the work technically sound?

Yes

Are sufficient details of methods and materials provided to allow replication by others?

Yes

Are the datasets clearly presented in a useable and accessible format?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Conservation Genomics, Ecology and Evolution

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 29 April 2022

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Petr Kotlik 🗓

Laboratory of Molecular Ecology, Institute of Animal Physiology and Genetics of the Czech Academy of Sciences, Liběchov, Czech Republic

This is a short report presenting an annotated complete genome assembly for the water vole, a small mammal species that is widespread in continental Europe but declining and endangered in Britain. The high-quality genome presented here will be an important resource for studies addressing conservation genomics and other questions about the biology of the water vole.

The manuscript is clearly presented. I have not found any problems in it and therefore have no suggestions for changes.

Is the rationale for creating the dataset(s) clearly described?

Yes

Are the protocols appropriate and is the work technically sound?

Yes

Are sufficient details of methods and materials provided to allow replication by others?

Yes

Are the datasets clearly presented in a useable and accessible format?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Evolutionary biology, population genomics, zoology, vertebrates

I confirm that I have read this submission and believe that I have an appropriate level of

expertise to confir	m that it is of an a	acceptable scienti	fic standard.	