GigaScience Draft genome of the Reindeer (Rangifer tarandus) --Manuscript Draft--

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Abstract:	Background: Reindeer (Rangifer tarandus) is the only fully domesticated species in the Cervidae family, and is the only cervid with a circumpolar distribution. Unlike all other cervids, female reindeer regularly grow cranial appendages (antlers, the defining characteristics of cervids), as well as males. Moreover, reindeer milk contains more protein and less lactose than bovid milk. A high quality reference genome of this species will assist efforts to elucidate these and other important features in the reindeer. Findings: We obtained 723.2 Gb (Gigabase) of raw reads by an Illumina Hiseq 4000 platform, and a 2.64 Gb final assembly, representing 95.7% of the estimated genome (2.76 Gb according to k-mer analysis), including 92.6% of expected genes according to BUSCO analysis. The contig N50 and scaffold N50 sizes were 89.7 kilo base (kb) and 0.94 mega base (Mb), respectively. We annotated 21,555 protein-coding genes and 1.07 Gb of repetitive sequences by de novo and homology-based prediction. Homology-based searches detected 159 rRNA, 547 miRNA, 1,339 snRNA and 863 tRNA sequences in the genome of R. tarandus. The divergence time between R. tarandus, and ancestors of Bos taurus and Capra hircus, is estimated to be 29.55 million years ago (Mya). Conclusions: Our results provide the first high-quality reference genome for the reindeer, and a valuable resource for studying evolution, domestication and other unusual characteristics of the reindeer		
Corresponding Author:	Wen Wang		
Corresponding Author Secondary Information:			
Corresponding Author's Institution:			
Corresponding Author's Secondary Institution:			
First Author:	Zhipeng Li		
First Author Secondary Information:			
Order of Authors:	Zhipeng Li		
	Zeshan Lin		
	Lei Chen		
	Hengxing Ba		
	Yongzhi Yang		
	Kun Wang		
	Qiang Qiu		
	Guangyu Li		
	Wen Wang		

Order of Authors Secondary Information:	
Opposed Reviewers:	Glenn Yannic, PhD Associate Professor glenn.yannic@univ-smb.fr
	Steeve Cote, Dr Steeve.Cote@bio.ulaval.ca
	Louis Bernatchez Iouis.bernatchez@bio.ulaval.ca
	Knut Roed knut.roed@nmbu.no
	Juha Kantanen, Prof juha.kantanen@luke.fi
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3 4 5 6	1	Draft genome of the Reindeer (Rangifer tarandus)
7 8	2	Zhipeng Li ^{1*} , Zeshan Lin ^{2*} , Lei Chen ^{2*} , Hengxing Ba ^{1*} , Yongzhi Yang ² , Kun
9 10 11	3	Wang ² , Wen Wang ^{2#} , Qiu Qiang ^{2#} , Guangyu Li ^{1#}
13 14	4	¹ Jilin Provincial Key Laboratory for Molecular Biology of Special Economic
15 16 17	5	Animals, Institute of Special Animal and Plant Sciences, Chinese Academy of
18 19 20	6	Agricultural Sciences, Changchun, 130112, China
21 22	7	² Center for Ecological and Environmental Sciences, Northwestern Polytechnical
23 24 25	8	University, Xi'an 710072, China
26 27 28	9	[*] These authors contributed equally to this work.
29 30 31	10	[#] Corresponding authors: tcslgy@126.com (GL), qiuqiang@lzu.edu.cn (QQ),
32 33 34	11	wwang@wangwen-lab.org or wwang@mail.kiz.ac.cn (WW)
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13 Abstract

14	Background: Reindeer (Rangifer tarandus) is the only fully domesticated species in
15	the Cervidae family, and is the only cervid with a circumpolar distribution. Unlike all
16	other cervids, female reindeer regularly grow cranial appendages (antlers, the defining
17	characteristics of cervids), as well as males. Moreover, reindeer milk contains more
18	protein and less lactose than bovids' milk. A high quality reference genome of this
19	specie will assist efforts to elucidate these and other important features in the reindeer.
20	Findings: We obtained 723.2 Gb (Gigabase) of raw reads by an Illumina Hiseq 4000
21	platform, and a 2.64 Gb final assembly, representing 95.7% of the estimated genome
22	(2.76 Gb according to k-mer analysis), including 92.6% of expected genes according
23	to BUSCO analysis. The contig N50 and scaffold N50 sizes were 89.7 kilo base (kb)
24	and 0.94 mega base (Mb), respectively. We annotated 21,555 protein-coding genes
25	and 1.07 Gb of repetitive sequences by de novo and homology-based prediction.
26	Homology-based searches detected 159 rRNA, 547 miRNA, 1,339 snRNA and 863
27	tRNA sequences in the genome of R. tarandus. The divergence time between R.
28	tarandus, and ancestors of Bos taurus and Capra hircus, is estimated to be 29.55
29	million years ago (Mya).
30	Conclusions: Our results provide the first high-quality reference genome for the
31	reindeer, and a valuable resource for studying evolution, domestication and other

- 32 unusual characteristics of the reindeer.

 33 Keywords: *Rangier tarandus*, whole genome sequencing, assembly, annotation

34 Background information

The Cervidae is the second largest family in the suborder Ruminantia of the Artiodactyla, which are distributed across much of the globe in diverse habitats, from arctic tundra to tropical forests [1, 2]. Interestingly, reindeer (*Rangifer tarandus*) is the only species with a circumpolar distribution (present in boreal, tundra, subarctic, arctic and mountainous regions of northern Asia, North America and Europe). It is also the only cervid having been fully domesticated, although some other species, such as the sika deer (Cervus nippon), which has been semi-domesticated for more than 200 years and still has strong wild nature. Antlers, male secondary sexual appendage, are the defining characteristic of cervids, which shed and regrow each year throughout an animal's life. However, reindeer do not follow this rule, with the exception in which females also bear shedding antlers. Moreover, reindeer milk contains greater amount of proteins, and lower amount of lactose compared to that of bovids [3]. Here, we report a high-quality reindeer reference genome using material from a Chinese individual, which will be useful in elucidating special characteristics of special cervid.

50 Data description

51 Animal and sample collecting

52 Fresh blood was collected from a two-year-old, female reindeer of a 53 domesticated herd maintained by Ewenki hunter-herders in the Greater Khingan Mountains, Inner Mongolia Autonomous Region, China (50.77° N, 121.47° E). The sample was immediately placed in liquid nitrogen, and was then stored at -80°C for later analysis.

57 Library construction, sequencing and filtering

Genomic DNA was extracted from the fresh blood. The isolated genomic DNA was then used to construct five short-insert libraries (200, 250, 350, 400 and 450 base pair, bp) and four long-insert libraries (3, 6.5, 11.5 and 16 kb) following standard protocols provided by Illumina. Then, 150 bp paired-end sequencing was performed to generate 723.2 Gb raw data, using a whole genome shotgun sequencing strategy on an Illumina Hiseq 4000 platform (Table S1). To improve the quality of reads, we trimmed low-quality bases from both sides of reads and removed reads with more than 5% of uncalled ("N") bases. Then reads of all libraries were corrected by SOAPec (version 2.03) [4]. Finally, clean reads amounting to 615 Gb were obtained for genome assembly.

68 Evaluation of genome size

The estimated genome size is 2.76 Gb according to k-mer analysis, based on the
following formula: G = k-mer_number/k-mer_depth (Figure S1) [5]. All the clean
reads provide approximately ~ 220-fold mean coverage.

72 Genome assembly

We used SOAPdenovo (version 2.04) with optimized parameters (pregraph -K 79 – d 0; map -k 79; scaff -L 200) to construct contigs and original scaffolds [5]. All reads were aligned onto contigs for scaffold construction by utilizing the paired-end information. Gaps were filled using reads from three libraries (200, 250 and 350 bp) with GapCloser (version 1.12) [6]. The final reindeer genome assembly is 2.64 Gb long, including 95.7 Mb (3.6%) of unknown bases, smaller than that of the domestic goat (*Capra hircus*, 2.92 Gb) [7], and similar to that of sheep (*Ovis aries*, 2.61 Gb) [8]. The contig N50 (> 200 bp) and scaffold N50 (> 500 bp) sizes are 89.7 kb and 0.94 Mb, respectively (Table 1).

82 Quality assessments of the assembled genome

We used BUSCO (benchmarking universal single-copy orthologs, version 2.0) software to assess the genome completeness [9]. Our assembly covered 92.6% of the core genes, with 3,803 genes being complete (Table S2). Feature-response curve (FRC, version 1.3.1) method [10] was then used to evaluate the trade-off between the assembly's contiguity and correctness. The results indicate that it has similar accumulated curve compare to published high quality assemblies for ruminant genomes including cattle, goat, and sheep (Figure S2). Subsequently, synteny analysis was applied to identify differences between the assembled genome and the domestic goat (*Capra hircus*) genome (Figure S3). 83.95% of two genome sequences could be 1:1 aligned, the average nuclear distance (percentage of different base pairs in the syntenic regions) was 7.18% (Figure S4). In addition, the density of different

94 types of break points (edges of structural variation) are about 69.88 per Mb (Table
95 S3). These results suggest that the reindeer genome assembly is of good level of
96 contiguity and correctness.

Genome annotation

To annotate the reindeer genome we initially used LTR FINDER [11] and RepeatModeller (version 1.0.4; http://www.repeatmasker.org/RepeatModeler.html) to find repeats. Next, RepeatMasker (version 4.0.5) [12] was used (with -nolow -no_is -norna -parallel 1 parameters) to search for known and novel transposable elements (TE) by mapping sequences against the *de novo* repeat library and Repbase TE library (version 16.02) [13]. Subsequently, tandem repeats were annotated using Tandem Repeat Finder (version 4.07b; with 2 7 7 80 10 50 2000 -d -h parameters) [14]. In addition, we used RepeatProteinMask software [12] with -no LowSimple -p value 0.0001 parameters to identify TE-relevant proteins. The combined results indicate that repeat sequences cover about 1.07 Gb, accounting for 40.4% of the reindeer genome assembly (Table S4).

109 The rest of the reindeer genome assembly was annotated using both *de novo* and 110 homology-based gene prediction approaches. For *de novo* gene prediction, we utilized 111 SNAP (version 2006-07-28), GenScan [15], glimmerHMM and Augustus (version 112 2.5.5) [16] to analyze the repeat-masked genome. For homology-based predictions, 113 sequences encoding homologous proteins of *Bos taurus* (Ensemble 87 release), *Ovis* 114 *aries* (Ensemble 87 release) and *Homo sapiens* (Ensemble 87 release), were aligned to the reindeer genome using TblastN (version 2.2.26) with an (E)-value cutoff of 1 e-5. Genwise (version wise2.2.0) [17] was then used to annotate structures of the genes. The *de novo* and homology gene sets were merged to form a comprehensive, non-redundant gene set using EVidenceModeler software (EVM, version 1.1.1), which resulted in 21,555 protein-coding genes (Table S5). We then compared the reindeer genome with species which used in homology prediction, and there is no significant difference among the four species in gene length and exon length distribution (Figure S5).

Next, we searched the KEGG, TrEMBL and SwissProt databases for best matches to the protein sequences yielded by EVM software, using BLASTP (version 2.2.26) with an (E)-value cutoff of 1 e-5, and searched Pfam, PRINTS, ProDom and SMART databases for known motifs and domains in our sequences using InterProScan software (version 5.18-57.0). At least one function was assigned to 19,004 (88.17%) of the detected reindeer genes through these procedures (Table S6). The reads from short-insert length libraries then were mapped to the reindeer genome with BWA (version 0.7.12-r1039) [18], then called single nucleotide variant (SNV) by SAMtools (version 1.3.1) [19]. Finally, we performed SnpEff (version 4.30) [20] to identify the distribution of SNV in the reindeer genome (Table S7).

In addition, we predicted rRNA-coding sequences based on homology with human rRNAs using BLASTN with default parameters. To annotate miRNA and snRNA genes we searched the Rfam database (release 9.1) with Infernal (version

б

0.81), and annotated tRNAs using tRNAscan-SE (version 1.3.1) software with default
parameters. The final results identified 159 rRNAs, 547 miRNAs, 1,339 snRNAs and
863 tRNAs (Table S8).

139 Species-specific genes and phylogenetic relationship

We clustered the detected reindeer genes in families by using OrthoMCL [21] with an (E)-value cutoff of 1 e-5, and a Markov Chain Clustering with default inflation parameter in an all-to-all BLASTP analysis of entries for five species (Homo sapiens, Equus caballus, Capra hircus, Bos taurus, and Rangifer tarandus). The result showed that 335 gene families were specific to the reindeer (Figure S6). Moreover, we identified 7,505 single-copy gene families from these species and aligned coding sequences in the families using PRANK (version 3.8.31) [22]. Subsequently, 4D-sites (four-fold degenerated sites) were extracted to construct a phylogenetic tree by RAxML (version 7.2.8) [23] with GTR+G+I model. Finally, phylogenetic analysis using PAML MCMCtree (version 4.5) [24], calibrated with timings published of the divergence of the reference species (http://www.timetree.org/), indicated that Rangifer tarandus, Bos taurus and Capra hircus diverged from a common ancestor approximately 29.6 (25.4-31.7) Mya (Figure S7).

154 Conclusion

In summary, we report the first sequencing, assembly and annotation of the

reindeer genome, which will be useful in analysis of the genetic basis of the unique characteristics of reindeer, and broader studies on ruminants. Availability of supporting data The raw data have been deposited in Genome Sequence Archive (GSA), under BIG Data Center, Beijing Institute Genomics (BIG), Chinese Academy of Science, with the project accession PRJCA000451. Abbreviations Gb: giga base; bp: base pair; kb: kilo base; Mb: mega base; TE: transposable

164 element; EVM: EVidenceModeler; BUSCO: benchmarking universal single-copy
165 orthologs; FRC: feature-response curves; SNV: single nucleotide variant; Mya:
166 million years ago

167 Acknowledgements

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174 Competing interests

175 The authors declare that they have no competing interests.

176 Authors' contributions

- 177 ZPL collected the samples; ZSL, CL ZPL, YZ, KW and HB analyzed the data;
- 178 ZSL, QQ and ZPL wrote the manuscript; GL, ZL, QQ and WW conceived the study.

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Tables

Туре	Scaffold (bp)	Contig (bp)
Total number	58,765	117,102
Total length	2,832,785,815	2,732,476,387
N50 length	986,392	91,805
N90 length	151,297	17,480
Max length	4,664,725	770,474
GC content(%)	41.24	40.98

247 Table 1 Summary of the genome assembly of *Rangier tarandus*

Figure legends

on four-fold degenerated sites. Estimated divergence times are shown above the

250 nodes. MYA, million years ago.





Figure 1. Phylogenetic relationships of *Rangier tarandus* and four species based on four-fold degenerated sites. Estimated divergence times are shown above the nodes. MYA, million years ago.

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