Data Article

Draft genome sequence of Japanese wood mouse, Apodemus speciosus

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A R T I C L E  I N F O

Article history:
Received 26 August 2017
Received in revised form 7 October 2017
Accepted 26 October 2017
Available online 31 October 2017

Keywords:
Rodent
Mouse
Apodemus speciosus
Phylogeography
Mammal
Evolution

A B S T R A C T

The wood mouse (genus Apodemus) is one of the most common rodents in broad-leaf forests in the temperate zone of the Palaearctic region. Molecular studies of wood mice have critically enhanced the understanding of their evolution and ancestral biogeographic events. However, their molecular data are currently only limited to partial mitochondrial sequences and a few genes. Therefore, we sequenced the wood mouse genome to facilitate the acquisition of useful resources for inferring their molecular evolution. We sampled a wild wood mouse at Tsukuba, Japan, and sequenced its whole-genome using the Illumina Hiseq. 2000. To reduce the risk of non-randomness, three paired-end libraries (insert sizes: 150, 300, and 500 bp) and, two mate-pair reads (insert sizes: 8 and 20 kbp) were constructed. In total, we generated approximately 210 Gbp data. From these sequences, we reconstructed 336,124 scaffolds. These data will enhance our understanding of the evolution and ecological factors that affect their genetic constitution. The genome scaffolds generated are
available in the National Center Biotechnology Information (NCBI) BioProject with accession number PRJDB5914.
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| Specifications Table |
|-----------------------|
| **Subject area**      | biology                                     |
| **More specific sub-**| Genomics                                    |
| **ject area**          |                                             |
| **Type of data**       | Genome sequences                            |
| **How data was**       | High throughput DNA sequencing using Hiseq. 2000 |
| **acquired**           |                                             |
| **Data format**        | FASTA format                                |
| **Experimental**       |                                             |
| **factors**            | Total DNA extraction from liver samples     |
| **Experimental**       |                                             |
| **features**           | Freshly collected healthy liver was used for total DNA extraction |
| **Data source**        |                                             |
| **location**           | Tsukuba, Ibaraki, Japan                     |
| **Data accessibility** | The genome scaffolds generated are available in the National Center Biotechnology Information (NCBI) BioProject with accession number PRJDB5914. |
| **Related research**   |                                             |
| **article**            | Not applicable                              |

**Value of the Data**

- The Japanese wood mouse could be a model mammal for investigating the natural histories of organisms inhabiting the Japanese Archipelago.
- We sequenced the Japanese wood mouse genome to facilitate the acquisition of useful resources for inferring the molecular evolution of this species.
- This data allows other researchers focusing on this species to start genome-wide analysis.

1. Data

The genus *Apodemus*, which is a member of the subfamily Murinae, is one of the most species rich taxa of the murine rodents, next to *Rattus* and *Mus*, which are the first and second speciose genera, respectively [1]. Species belonging to *Apodemus*, known as field or wood mice, are distributed in the temperate region of the Eurasian continent including the Japanese Archipelago. According to previous molecular phylogenetic analyses, *Apodemus* diversified in Asia and Europe, around the early Pliocene and early Pleistocene eras, respectively [2]. Deciphering the genome sequences of this genus would facilitate our understanding of the evolutionary trait of the group adapted to the temperate environment, with the aid of those of *Rattus* and *Mus*. The genome of a representative species of *Apodemus* from Europe, *Apodemus sylvaticus*, was sequenced by a research group at the Liverpool University according to the JGI GOLD database (https://gold.jgi.doe.gov; project ID = Gp0143296), but its project status is “permanent draft.” In Asian species, no genome sequence data are currently available.
The large Japanese wood mouse, *Apodemus speciosus*, is widely distributed in Japan from Hokkaido to Kyushu, including the remote peripheral islands. This species inhabits forests and grasslands, eats insects and acorns, and shows seasonal breeding in the wild. A recent phylogeographic analysis revealed that this species experienced a rapid population growth in response to the effect of the Quaternary ice ages [3]. Thus, the species could be a model mammal for investigating the natural histories of organisms inhabiting the Japanese Archipelago [4]. We sequenced the Japanese wood mouse genome to facilitate the acquisition of useful resources for inferring the molecular evolution of this species.

The genome was assembled using the ALLPATHS-LG program using two-step processes of assembling and scaffolding. The paired-end reads were first assembled into contigs, which were then joined to form concatenated scaffolds by using mate-pair reads. We reconstructed 647,983 contigs (N50 is 3.5 kbp). These contigs were joined in 336,124 scaffolds (Table 2), and the N50 value of the scaffolds was 47 kbp. The wood mouse genome sequenced is a potentially useful resource for the future molecular evolutionary analysis of rodents.

2. Experimental design, materials, and methods

2.1. Sample preparation and sequencing

We sampled the *A. speciosus* at Tsukuba, Japan (NIES ID; IB14-021, male). All rodent experiments were conducted in accordance with the guidelines for using wild mammals of the Mammal Society of Japan [5] and the rules of the National Institute for Environmental Studies for analysis and experimentation with environmental samples contaminated with radioactive materials. Genomic DNA was extracted from the liver using proteinase K and phenol:chloroform:isoamyl alcohol [6]. The library preparations and sequencings were conducted by the Eurofins Sequencing Service (Eurofins MWG Operon, Germany). Paired-end sequencing of each 100-bp end of the DNA fragments was performed using the Hiseq. 2000 sequencing system (Illumina, San Diego, CA, USA). We sequenced libraries showing five different insert sizes: 150 bp, 300 bp, 500 bp, 8 kbp, and 20 kbp.

| Insert size (bp) | No. of read pairs | No. of reads | Read length (bp) | Total data (GB) |
|-----------------|-------------------|--------------|-----------------|-----------------|
| 150             | 362,107,447       | 724,214,894  | 100             | 72.4            |
| 300             | 336,419,570       | 892,839,140  | 100             | 89.3            |
| 500             | 369,663,105       | 739,326,210  | 100             | 73.9            |
| 8000            | 22,370,797        | 44,741,594   | 100             | 4.5             |
| 20,000          | 12,377,116        | 24,754,232   | 100             | 2.5             |

Table 2
General features of the *Apodemus speciosus* genome.

| Feature                                      | Value            |
|----------------------------------------------|------------------|
| Number of contigs                            | 647,983          |
| Number of contigs per megabase (Mb)          | 189              |
| Number of scaffolds                          | 336,124          |
| Total contig length                          | 2,009,551,775    |
| Total scaffold length, with gaps             | 3,428,181,270    |
| N50 contig size (kb)                         | 3.5              |
| N50 scaffold size (kb)                       | 25               |
| N50 scaffold size (kb), with gaps            | 47               |
| Number of scaffolds per megabase (Mb)        | 98.05            |
| Median size of gaps in scaffolds             | 2846             |
| Median deviation of gaps in scaffolds         | 852              |
2.2. De novo genome assembly

We sequenced three paired-end libraries (insert sizes: 150, 300, and 500 bp) and two mate-pair reads (insert sizes: 8, and 20 kbp). In total, we generated approximately 210 Gbp data. We obtained 362107447, 336419570, 369663105, 22370797, and 12377116 reads for each library showing 150-bp, 300-bp, 500-bp, 8-kbp, and 20-kbp insert sizes, respectively (Table 1). The sequenced reads were used for de novo genome assembly using the ALLPATHS-LG program ver. 51927 under default parameter settings [7].

Acknowledgements

We would like to appreciate Dr. Hiroko Ishiniwa for the helpful discussions. This work was supported by an NIG Collaborative Research Program Grant to H. S. (No. 2015-A1-14) and a KAKENHI Grant-in-Aid for Young Scientists (B) to M.M. (No. 16K18613). The computational analysis utilized the supercomputer system of the National Institute of Genetics, Research Organization of Information and Systems.

Transparency document. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.dib.2017.10.063.

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