A Chromosome-Level Genome Assembly for the Rock Ptarmigan (Lagopus muta)

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Abstract:

The Rock Ptarmigan (*Lagopus muta*) is a cold-adapted, largely sedentary, game bird with a Holarctic distribution. The species represents an important example of an organism likely to be affected by ongoing climatic shifts across a disparate range. We provide here a high-quality reference genome and mitogenome for the Rock Ptarmigan assembled from PacBio HiFi and Hi-C sequencing of a female bird from Iceland. The total size of the genome is 1.03 Gb with a scaffold N50 of 71.23 Mb and a contig N50 of 17.91 Mb. The final scaffolds represent all 40 predicted chromosomes, and the mitochondria with a BUSCO score of 98.6%. Gene annotation resulted in 16,078 protein-coding genes out of a total 19,831 predicted (81.08% excluding pseudogenes). The genome included 21.07% repeat sequences, and the average length of genes, exons, and introns were, 33605, 394, and 4265 bp respectively. The availability of a new reference-quality genome will contribute to understanding the Rock Ptarmigan's unique evolutionary history, vulnerability to climate change, and demographic trajectories around the globe. As a wild relative of domesticated Galliformes, findings of avicultural importance may also be revealed.

Kev words: Reference Genome, *Lagopus muta*, Rock Ptarmigan, Hi-C

Introduction:

The Rock Ptarmigan (*Lagopus muta*, Montin 1776) is a grouse species with a wide distribution across the arctic and subarctic northern hemisphere. It has seasonally variable plumage ranging from almost entirely white in the winter to heavily mottled grey, rust, and brown in the breeding months (see fig. 1). Birds of the genus *Lagopus* are notable for having feathered legs and feet which likely serve to insulate them in cold habitats. The Rock Ptarmigan can be considered as a ring species with variable genetic diversity across its circumpolar range (Sahlman et. al., 2009, Kozma et. al., 2019; see fig. 2). Accordingly, Rock Ptarmigan are expected to be at long-term risk across much of their range due to

ongoing climatic changes and limited suitable habitat (Costanzi and Steifetten, 2019; Masanobu et al., 2019).

With the expected declines in cold specialist species as global temperatures rise (Chamberlain et. al., 2012; Scheffers et al. 2016; Scridel et al., 2019; Hoglund et al., 2021), resident birds such as the individual we used to generate the reference genome are particularly valuable to science as they are likely to display many special adaptations necessary for life in the arctic or at high altitude. Some populations of Rock Ptarmigan are considered near-threatened or endangered due to long-term population loss and habitat declines (Nakamura, 2014; Icelandic Institute of Natural History, 2018). The risks associated with declining genetic quality and environmental changes are not well understood, but might be better assessed with a locally sourced reference genome. For populations with robust historical demographics such as the Icelandic population (Gardarsson, 1988; Nielsen et al., 1986-2011), a locally sourced reference genome is exponentially more valuable in assessing demographic history.

The species nearest relatives include other members of Tetraonidae (order Galliformes). The mitochondrial genome of Rock Ptarmigan was previously made available along with the mitochondrial DNA of a sister species Willow Grouse (*Lagopus lagopus*; Sveinsdóttir and Magnússon, 2017). The Willow Grouse and Rock Ptarmigan are believed to have diverged as recently as 1 million years ago (Kozma et al., 2019) and often studied together as sister species. The white-tailed ptarmigan (*Lagopus leucura*) is the most closely related species with whole genome data available, having a common ancestor with other *Lagopus* taxa no older than 3 million years ago, although the genome assembly is not currently annotated (Kozma et al., 2019; Clark et. al., 2016; GenBank: GCA_019238085.1).

Here, we describe the first reference-quality genome assembly and annotations for Rock Ptarmigan. A combination of long-read and conformation capture sequencing technologies was used to assemble the 1.03 Gb haploid genome.

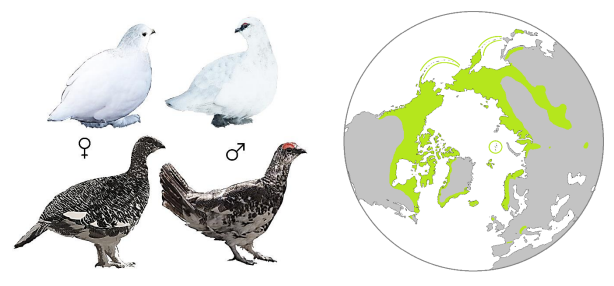


Figure 1

Diagram of the sexually dimorphic seasonal molt patterns of adult Rock Ptarmigan showing white winter plumage and mottled breeding colors.

Figure 2

A range map showing the global distribution of Rock Ptarmigan above 30° north.

Materials and methods:

Sample collection and PCR preparation:

High molecular weight DNA extraction was performed in laboratories at Uppsala University using special procedures associated with the DNeasy Blood & Tissue kits (Qiagen, Germany). As basis for the reference genome assembly and annotation, blood from a single female bird collected (shot) in Húsavík, northern Iceland, in 2018 was used (NCBI BioSample SAMN25144835) while additional blood, heart, muscle, brain, kidney, liver, ovaries, and spleen from a second bird was collected for RNA-seq to aid in gene prediction (NCBI BioSample SAMN26436951, SAMN29421920, SAMN29421921, SAMN29421922, SAMN29421923, SAMN29421924, SAMN29421925, and SAMN29421926 respectively). Materials from the birds used remain accessioned with the Icelandic Institute of Natural History in Garðabær, Iceland.

Sequencing:

Sequencing was achieved using two PacBio SMRT smart cells run on a Sequel II system, while Dovetail Genomics Hi-C Kits were processed on an Illumina NovaSeq 6000.

Genome assembly:

The genome was assembled following the Vertebrate Genome Project (VGP; Rhie, 2021) assembly pipeline. First, a kmer database was generated using Meryl (v.1.3) from the PacBio HiFi reads for reference-free genome evaluation and downstream assembly QC. The kmer size was set to 21 after running the best_k.sh script for the expected genome size (~1Gb) in Merqury (v.1.3; Rhie et al., 2020). PacBio HiFi reads were assembled using hifiasm (v. 0.15.1-r334; Cheng et al., 2021), followed by a round of purge_dups (v. 1.2.5; Guan et al., 2020) incorporating minimap2 (v.2.17-r941). Each of the previous steps was followed by assembly evaluation which included contig/scaffold statistics calculated via Assembly_stats (v. 0.1.4; 10.5281/zenodo.3968775), BUSCO (v. 5.3.1), and kmer spectrum plots using Merqury (v.1.3; Rhie et al., 2020). The assembly was scaffolded using the Hi-C reads. Briefly, reads were first aligned to the assembly using the VGP modified version of the Arima mapping pipeline that uses bwa mem (v.0.7.17-r1188) and samtools (v. 1.19) for alignment and Picard (v.2.10.3) for 5′-end filtering and duplication removal. Scaffolding was performed using Salsa2 (v.2.3) and evaluated using BUSCO and scaffold statistics.

The Hi-C reads were then mapped back to the scaffolded assembly using the same pipeline as in the previous step and the resulting bam file was converted to pretext format using PretextMap (v. 0.1.7). The resulting Hi-C contact maps were visualized and edited in PretextView (v. 0.2.5). The assembly then was curated by the Wellcome Sanger Institute.

The mitochondrial genome was assembled separately from both raw reads and contigs using MitoHifi (v. 2.2; Uliano-Silva et. al., 2021) with automatic alignment to the Japanese Rock Ptarmigan (*L. muta japonica*; Yonezawa and Nishibori, 2020) via built-in features from the MitoFinder dependency (v. 1.4.1; Allio et. al., 2020).

The completed genome assembly is publicly available in NCBI under accession number GCA_023343835.1.

Genome annotation:

The Rock Ptarmigan reference genome was annotated using the standard NCBI Eukaryotic Genome Annotation Pipeline version 10.0. A detailed summary of the pipeline is available online at: https://www.ncbi.nlm.nih.gov/genome/annotation_euk/process/. In contrast to previous iterations, this version of the pipeline used RFAM (v.14.6; Kalvari et al., 2021) for discovery of small non-coding RNA's and STAR (Dobin et al. 2013) for alignment of RNA-seq reads from our supplementary tissues. The pipeline has stable use of several tools including BUSCO (v. 4.1.4; Manni et al., 2021) and Splign (Kapustin et al., 2008) among others.

For calculation of genomic masking, the Rock Ptarmigan had its genomic sequence masked by WindowMasker (Morgulis et al., 2006). Annotation of the mitochondrial genome was achieved via manual comparison with the extant published Icelandic Rock Ptarmigan genome in addition to automatic annotation using MITOS WebServer (Bernt et. al., 2013).

Results:

Sequencing and Assembly Results:

The final assembly sequence is 1,026,771,810 base pairs long, with 71,937 gap bases (0.007%). The genome assembly includes 375 contigs arranged on 165 scaffolds. The scaffold N50 is 71,229,700 bp with an L50 of 5. The Contig N50 is 17,905,263 bp with an L50 of 19.

Average coverage across the genome is 57.75x. In total 38 autosomes were identified, with 18 unlocalized sequences among them. Additional W and Z allosomes were described with only a single unlocalized sequence found on the W. Assembly summary statistics appear significantly better than the current *Gallus gallus* reference genome (GRCg6a), and are modest in comparison to the most recently annotated *Gallus gallus* individual (bGalGal1.mat.broiler.GRCg7b; See Table 1 below).

Statistical analysis to determine completeness and quasi-variance showed expected values (See Table 2 below)

| | Lagopus muta Reference Genome (bLagMut1) | Gallus gallus Most Recent Annotation (bGalGal1.mat.broiler.GRCg7b) | Gallus gallus Reference Genome (GRCg6a) |
|--------------------------|--|---|---|
| Total length | 1,026,771,810 | 1,053,332,251 | 1,065,348,650 |
| Total ungapped length | 1,026,699,873 | 1,049,948,333 | 1,055,564,190 |
| Gaps between scaffolds | 0 | 0 | 68 |
| Number of scaf- folds | 165 | 214 | 524 |
| Scaffold N50 | 71,229,700 | 90,861,225 | 20,785,086 |
| Scaffold L50 | 5 | 4 | 12 |
| Number of contigs | 375 | 677 | 1,402 |
| Contig N50 | 17,905,263 | 18,834,961 | 17,655,422 |
| Contig L50 | 19 | 18 | 19 |
| Chromosomes and plasmids | 41 | 42 | 34 |
| Component sequences | 165 | 677 | 2,243 |

Table 1

A series of "global" statistics published in the public release of the Rock Ptarmigan reference genome on NCBI indicating the completeness of the new reference genome in comparison to the gold standard Chicken reference genome and the most recently annotated Chicken reference genome.

| | Completeness | | Quasi-variance | | | | |
|--------------|--------------|-----------|----------------|-------|----------|---------|----------|
| bLagMut1.pri | 934494250 | 992616058 | 94.1446 | 10953 | 1.03E+09 | 62.9412 | 5.08E-07 |
| bLagMut1.alt | 856101045 | 992616058 | 86.2469 | 7185 | 9.23E+08 | 64.3112 | 3.71E-07 |
| both | 989631110 | 992616058 | 99.6993 | 18138 | 1.95E+09 | 63.5365 | 4.43E-07 |

Table 2

A series of statistics showing completeness and quasi-variance scores for the bLagMut1 assembly.

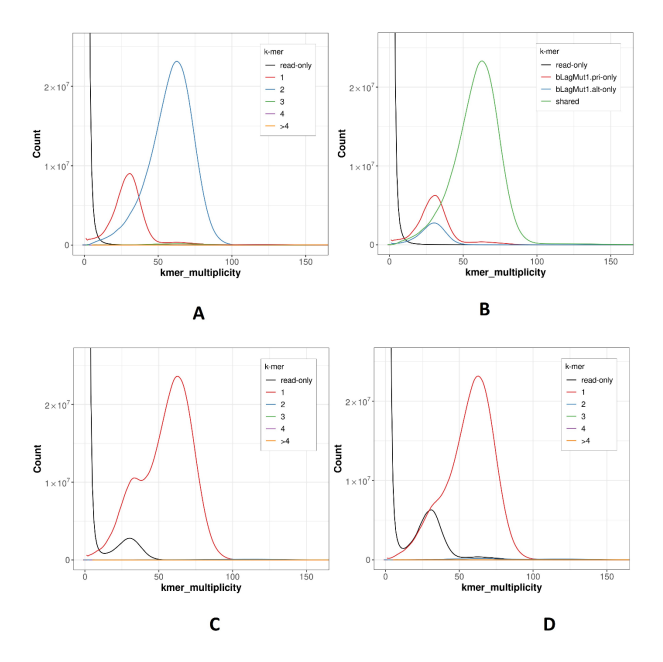


Figure 3

Outputs from Merqury showing kmer distribution according to: (A.) Spectra-cn plot of the bLagMut1 complete assembly, (B.) spectra-asm plot of the bLagMut1 complete assembly, (C.) spectra-cn plot of the bLagMut1 primary assembly, and (D.) spectra-cn plot of the bLagMut1 alternate assembly.

Genome Annotation:

In total 20,110 genes and pseudogenes were identified by combining gene prediction and similarity approaches, with approximately 80% identified as protein coding. The annotated genes showed a 98.6% completeness score against 98.9% for the whole genome when set against the BUSCO avian dataset (aves_odb10 lineage) and indicating 0.9% of genes missing from the annotated assembly. The annotation

and associate summary statistics are available in NCBI's RefSeq genome record for the reference (Pruitt et al., 2013). The contents of the report are summarized in Table 3.

| | Lagopus muta Reference Genome Annotation (bLagMut1) | Gallus gallus Most Recent Annotation (bGalGal1.mat.broiler.GRCg7b) |
|--------------------------|---|---|
| Genes and Pseudogenes | 20,110 | 25,635 |
| Protein Coding Genes | 16,078 | 18,023 |
| Non-Coding Genes | 3,738 | 7,330 |
| mRNA | 43,785 | 68,670 |
| IncRNA | 5,431 | 10,062 |
| tRNA | 306 | 303 |
| CDSs | 43,793 | 68,683 |
| Introns (mean length) | 206,142 (4,265) | 241,290 (4,145) |
| Exons (mean length) | 229,018 (394) | 262,919 (490) |
| Mean Gene Size | 33kb | 28kb |
| Maximum Gene Size | 1.6 Mb | 1.3 Mb |
| BUSCO Score | 98.6% | 98.7% |

Table 3

A comparative table showing the relative accuracy and completeness of the *Lagopus muta* reference annotation (NCBI *Lagopus muta* Annotation Release 100) against the most recently complete annotation of the chicken genome (NCBI *Gallus gallus* Annotation Release 106).

Mitochondrial Genome:

The mitochondrial DNA was described with all 13 expected protein coding sequences. The complete mitochondrial genome was assembled from reads using MitoHifi (Uliano-Silva et. al., 2021) and analyzed for accuracy through comparative analysis. With our addition, there are now four extant mitochondrial genomes published for the Rock Ptarmigan; Two from Iceland, one from Japan, and one from Siberia (Sveinsdóttir and Magnússon, 2017; Yonezawa et. al., 2020; Wang et. al., 2017). Using the ClustalW package embedded in BioEdit (Thompson et. al., 1994; Hall, 1999), we found a total of 24 bases divergent from the previously published Icelandic Rock Ptarmigan mitogenome in a manual review. Of these divergences 14 appeared in coding regions and 8 appeared unique to the previously published individual and our calls at those locations were conserved in the other Rock Ptarmigan populations. None of the polymorphisms observed between the populations appeared to be uniquely conserved in the

Icelandic Population. Analysis of pairwise distances using phylogenetic tree software in Mega11 (Tamura et. al., 2021) showed clear grouping of the Rock Ptarmigan separated from the Willow Ptarmigan, as previously reported (Sveinsdóttir and Magnússon, 2017).

Discussion:

Our new avian reference genome includes a highly complete set of information with 99.994% of the 1.03 Gb described matching to 40 haploid chromosomes and the mitochondrial genome. Other recent works have aimed to unlock the potential provided by Rock Ptarmigan genetics (Kozma, 2019; Sigmarsdóttir, 2022). In comparison to other recently published genomes, it is of clearly high quality.

Though Rock Ptarmigan has been globally identified as Least Concern by the IUCN in recent years, there have been regional fluctuations in its status and some nations identify the species as threatened due to long term declines (IUCN, 2022; European Commission, 2022; Icelandic Institute of Natural History, 2018). It is well established that Arctic species such as Rock Ptarmigan may be disproportionately affected by climate change with an expected poleward contraction of species ranges (Birdlife International, 2015). For more disparate populations such as those in the mountains of Honshu, the European Alps, and the Pyrenees, rising tree lines may entirely squeeze the Rock Ptarmigan out of its montane niches as has been suggested broadly for alpine habitats (Dirnböck et al., 2011; MRI EDW Working Group, 2015), and some closely related species (Jackson et al., 2016). Given the species' unique evolutionary adaptations to life in the arctic and uncertain future, this reference genome should be recognized as one with broad conservation implications.

Many wildlife species are difficult to study at the genomic level due to limited specimen availability and limitations on procurement (Hope et al., 2018; Kemp, 2015). Because the Rock Ptarmigan is a widespread game bird, it is particularly useful for genomic studies. Hunters have the potential to contribute robust data regarding the species trends and may continue to contribute both historical and new specimen materials for research (Cretois et al., 2020). Because of the species close cultural connection to some regions and history as a food source, the Rock Ptarmigan may benefit from additional conservation efforts from an involved public or concerned hunters and may be a good candidate for flagship status (McGowan et al., 2020).

Future studies into Rock Ptarmigan genomics may benefit from a history of captive breeding and the availability of nearly wild birds. Rock Ptarmigan has been utilized as food and traded since prehistory with human dispersal possible as early as the Viking age (McGovern et al., 2006), but in the last four decades of the modern era, birds from Svalbard have been regularly brought to a colony housed in Tromsø for research purposes (Stokkan, 1988). Already, Rock Ptarmigan hatched and raised in captivity have been used for gene expression studies to understand circadian rhythms and investigations of the cecal microbiome (Appenroth et al., 2021; Salgado-Flores et al., 2019). This represents a valuable opportunity for tailored future study.

Among avian diversity, the birds in the family Galliformes represent less than 3% of all species but have an outsized impact on global economics with Chickens, Turkeys, Pheasants, Quails, and Grouse all being regularly consumed. Among the available avian genomes (Bravo et al, 2021) those in order Galliformes are represented with 26 species assemblies currently available on NCBI (approximately 5% of all extant; Sayers et al., 2022). Among these, 68 assemblies have been completed and the domestic chicken has been assembled 30 times. This highlights a commercial implication for Rock Ptarmigans as they have many unique adaptations that could be of importance to the poultry industry.

Much work has been done using the chicken genome for past studies (Bert, 2005; Li et al., 2022) and given the usefulness of wild relatives for research into domesticated species (Li et al., 2020; Jackson et al., 2016) the Rock Ptarmigan may prove to be an exceedingly useful model for understanding other Galliformes. This relationship will surely have limitations in the genomic realm as more distantly related species are less informative at finer scales than those that are closely related (Scutari et al., 2016). However, if the Rock Ptarmigan's genes tailored to arctic landscapes can be used to better understand genetic pathways for cold weather survival or improved forage capabilities, viable improvements may be identified for commercially important species. Because the Rock Ptarmigan is more phylogenetically basal to many modern domesticated Galliformes, it may aid in the identification of gene developments responsible for ancestral traits.

Taking all of this into consideration, the arrival of a Rock Ptarmigan reference genome makes the species exceptionally well positioned for investigation across a broad range of scientific inquiry. With links to arctic/alpine biomes, hunting cultures, and the poultry industry, the Rock Ptarmigan reference genome provides a unique opportunity to capitalize on a species at the intersection of many issues of global significance.

Significance statement:

The Rock Ptarmigan is a widespread bird species of economic and nutritional importance to large portions of the northern hemisphere. Though small parts of the Rock Ptarmigan's genetic profile were previously understood, the massive effort undertaken to sequence and annotate the whole genome provides an ability to understand the species at a molecular level. This vertebrate genome allows for new critical assessment of the Rock Ptarmigan and related species at the individual, population, and the environmental level.

Conflicts of Interest:

We declare no conflicts of interest.

<u>Preprint Statement:</u> The authors wish to advise that the current preprint has been compiled, revised, and submitted, almost exclusively by Patrik Rödin-Mörch, Theodore E. Squires, and Giulio Formenti with minimal oversight from the other listed authors. While the other authors will be included in final published versions of this manuscript with reflections of their input on the contents, we can not at this time say that the contents of this paper, particularly in the discussion section, represent the views of all listed authors. Much deserved acknowledgements to the laboratory technicians who assisted with this project will also be added.

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Data Availability Statement (required):

The final annotation has been publicly released and uploaded according to the high standards of the Earth BioGenome Project (Lewin et al., 2018). The genome assembly, including the raw shotgun sequencing data, and the mitochondrial genome has been uploaded to NCBI and is available at https://www.ncbi.nlm.nih.gov/assembly/GCA_023343835.1; BioProject: PRJNA836583; BioSample: SAMN25144835

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