

GigaScience, 7, 2018, 1–14

doi: 10.1093/gigascience/giy044 Advance Access Publication Date: 2 May 2018 Research

RESEARCH

Whole-genome *de novo* sequencing reveals unique genes that contributed to the adaptive evolution of the Mikado pheasant

Chien-Yueh Lee^{1,†}, Ping-Han Hsieh^{1,†}, Li-Mei Chiang¹, Amrita Chattopadhyay², Kuan-Yi Li^{3,4}, Yi-Fang Lee¹, Tzu-Pin Lu⁵, Liang-Chuan Lai⁶, En-Chung Lin⁷, Hsinyu Lee^{1,8,9}, Shih-Torng Ding^{7,9}, Mong-Hsun Tsai^{2,9,10,11}, Chien-Yu Chen^{3,9,12,*} and Eric Y. Chuang ^{1,2,13,*}

¹Graduate Institute of Biomedical Electronics and Bioinformatics, National Taiwan University, Taipei 10617, Taiwan, ²Bioinformatics and Biostatistics Core, Center of Genomic Medicine, National Taiwan University, Taipei 10055, Taiwan, ³Department of Bio-Industrial Mechatronics Engineering, National Taiwan University, Taipei 10617, Taiwan, ⁴Institute of Plant and Microbial Biology, Academia Sinica, Taipei, 11529, Taiwan, ⁵Institute of Epidemiology and Preventive Medicine, National Taiwan University, Taipei 10055, Taiwan, ⁶Graduate Institute of Physiology, National Taiwan University, Taipei 10051, Taiwan, ⁷Department of Animal Science and Technology, National Taiwan University, Taipei 10617, Taiwan, ⁸Department of Life Science, National Taiwan University, Taipei 10672, Taiwan, ¹⁰Institute of Biotechnology, National Taiwan University, Taipei 10672, Taiwan, ¹¹Agricultural Biotechnology Research Center, Academia Sinica, Taipei 10672, Taiwan University, Taipei, Taiwan, ¹²Center for Systems Biology, National Taiwan University, Taipei 10672, Taiwan and ¹³Graduate Institute of Chinese Medical Science, China Medical University, Taichung 40402, Taiwan

*Correspondence address. Eric Y. Chuang, Graduate Institute of Biomedical Electronics and Bioinformatics, National Taiwan University, Taipei 10617, Taiwan. Tel: +886-2-3366-3660; Fax: +886-2-3366-3682; E-mail: chuangey@ntu.edu.tw; Chien-Yu Chen, Department of Bio-Industrial Mechatronics Engineering, National Taiwan University, Taipei 10617, Taiwan. Tel: +886-2-3366-5334; E-mail: chienyuchen@ntu.edu.tw [®] http://orcid.org/0000-0003-2530-0096 [†]These authors contributed equally to the work.

Abstract

Background: The Mikado pheasant (Syrmaticus mikado) is a nearly endangered species indigenous to high-altitude regions of Taiwan. This pheasant provides an opportunity to investigate evolutionary processes following geographic isolation. Currently, the genetic background and adaptive evolution of the Mikado pheasant remain unclear. **Results:** We present the draft genome of the Mikado pheasant, which consists of 1.04 Gb of DNA and 15,972 annotated protein-coding genes. The Mikado pheasant displays expansion and positive selection of genes related to features that contribute to its adaptive evolution, such as energy metabolism, oxygen transport, hemoglobin binding, radiation response, immune response, and

Received: 31 August 2017; Revised: 28 January 2018; Accepted: 13 April 2018

[©] The Author(s) 2018. Published by Oxford University Press. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

DNA repair. To investigate the molecular evolution of the major histocompatibility complex (MHC) across several avian species, 39 putative genes spanning 227 kb on a contiguous region were annotated and manually curated. The MHC loci of the pheasant revealed a high level of synteny, several rapidly evolving genes, and inverse regions compared to the same loci in the chicken. The complete mitochondrial genome was also sequenced, assembled, and compared against four long-tailed pheasants. The results from molecular clock analysis suggest that ancestors of the Mikado pheasant migrated from the north to Taiwan about 3.47 million years ago. **Conclusions:** This study provides a valuable genomic resource for the Mikado pheasant, insights into its adaptation to high altitude, and the evolutionary history of the genus *Symaticus*, which could potentially be useful for future studies that investigate molecular evolution, genomics, ecology, and immunogenetics.

Keywords: Mikado pheasant; Syrmaticus mikado; long-tailed pheasant; whole-genome sequencing; de novo genome assembly; adaptive evolution

Background

The Mikado pheasant (*Syrmaticus mikado*), which is a long-tailed pheasant indigenous to Taiwan, belongs to the family *Phasianidae* in the order Galliformes (Additional file 1: Fig. S1A, B). The Mikado pheasant is known to inhabit a variety of habitats in the mountainous regions of central and southern Taiwan at very high elevations ranging from 1,600 to 3,500 meters [1, 2]. The Mikado pheasant faced endangerment due to hunting pressure and habitat destruction [3, 4] until it became protected under the Wildlife Conservation Act. Currently, the International Union for Conservation of Nature Red List has classified the Mikado pheasant as a nearly threatened species, showing a decreasing trend in the overall population, with an estimate of approximately 15,000 mature birds. The rare and precious Mikado pheasant is a national icon in Taiwan and is depicted on its 1000 dollar banknote.

The *de novo* genome assembly of endangered species is an effective approach to identify genomic signatures associated with environmental adaptation and behavioral attributes [5, 6]. Genome resources can also provide great insights into effective population size, genetic defects, and deleterious mutations [7, 8]. Moreover, reconstruction of a phylogenetic tree can reveal genetic relationships and evolutionary history [9-11]. Together, they can lead to the conservation and rescue of endangered species.

The Mikado pheasant possesses ideal characteristics for evolutionary research because of its flightlessness and habitat isolation. It is one of five long-tailed pheasants in the Syrmaticus genus, which forms a monophyletic group [12]. Due to limited molecular data, very few studies have been conducted to investigate the phylogenetic relationships and divergence time of species within the genus. Moreover, the Mikado pheasant is mainly found in Yushan National Park [13], which has numerous extremely high mountains that exceed an altitude of 3,000 meters (Additional file 1: Fig. S2). As high altitudes are associated with extremely cold climates and lower oxygen concentrations, hypoxic stress may be observed in the pheasant. Considering its importance as a species facing endangerment, the present unavailability of genetic information regarding the Mikado pheasant motivated the *de novo* assembly of its genome, followed by a detailed study of its genetic background and subsequent adaptive evolution.

Here, we report the whole-genome assembly of the Mikado pheasant and provide insights into its adaptive mechanisms. This genome-wide study reveals the evolutionary adaptation of the Mikado pheasant to high altitudes, including changes in gene family size and/or molecular signatures of positive selection associated with energy metabolism, oxygen transport, hemoglobin binding, radiation response, immune response, and DNA repair. The estimated time of divergence among the five long-tailed pheasant species reconstructs the evolutionary history of the lineage and allows us to propose a hypothesis for the biogeographical speciation of the Mikado pheasant. Additionally, the manually curated major histocompatibility complex (MHC) gene loci of the Mikado pheasant display evidence for molecular evolution with a high level of synteny, mainly across inverse regions in gene blocks, and several rapidly evolving genes in comparison with the chicken.

Data Description

The details about sample collection, library construction, sequencing, assembly, gene prediction, and annotation can be found in the Materials and Methods section.

Results

Genome assembly and annotation

In total, 171.7 Gb of raw DNA sequence reads (Additional file 1: Table S1) were generated, resulting in an approximately 160fold sequencing coverage based on the 1.07 Gb genome size estimated by KmerGenie [14]. The contigs were built and assembled into a 1.04 Gb sequence of the draft genome. The N50 lengths of the contigs and scaffolds were 13.46 kb and 11.46 Mb, respectively. The overall GC content of the Mikado pheasant genome was 41.13%, which is similar to that of the chicken, duck, turkey, and zebra finch (Additional file 1: Fig. S3). The size of the longest assembled sequence was 50.28 Mb, and 928 scaffolds were longer than 10 kb. The basic statistics of both the contigs and scaffolds assembled using MaSuRCA [15] are shown in Table 1. The cumulative length plots (Additional file 1: Fig. S4A, B) and the Nx plot for the scaffolds (Additional file 1: Fig. S5) showed that most of the draft genome consisted of large scaffolds; though many short scaffolds were present, they only contributed a small portion of the genome size.

Before performing gene prediction and annotation, the interspersed and low complexity regions were first masked using RepeatMasker (RepeatMasker, RRID:SCR_012954) [16]. Approximately 8.91% of the sequences were identified as interspersed repeats, 1.32% of the sequences were identified as long tandem repeat elements, and overall 11.46% of the total bases were identified (Additional file 1: Table S2). After masking the repeats and extrinsic data, an *ab initio* gene prediction was performed using Augustus (Augustus: Gene Prediction, RRID:SCR_008417) [17], followed by EVidenceModeler [18]. The final gene models comprised 27,254 transcripts (proteins). Of the predicted proteins, 15,972 (58.6%) could be strictly aligned to the National Center for Biotechnology Information (NCBI) nonredundant (NR) pro-



Figure 1: A chromosome-level comparison of the Mikado pheasant and the chicken. (A) A syntenic map of the Mikado pheasant and chicken genomes. The x-axis specifies the chromosome position in the chicken, whereas the y-axis specifies the scaffold position in the Mikado pheasant. The red dots (or lines) indicate that the sequences were aligned in the same orientation, and the blue dots indicate an alignment with a reverse complement. (B) A chord diagram of scaffolds with a total length greater than 500 kb and an alignment length greater than 10 kb. The orange perimeters specify the chromosomes (chr) of the chicken, whereas the purple perimeters specify the scaffolds (sc) of the Mikado pheasant. The red links represent the sequences aligned in the same orientation, and the blue links represent an alignment with a reverse complement. Yellow arrows indicate the scaffolds that were fully aligned, and gray arrows indicate the multiple alignment.



Figure 2: Evolution of gene families among various animal species. A phylogenetic tree was reconstructed based on 5,287 single-copy orthologs of 10 species. The most recent common ancestor (MRCA) contains 18,220 gene families that were used to examine gene families with expansions or contractions. The numbers of gene families with significant expansions and contractions are shown in red and blue, respectively, at each branch. The divergence times and associated 95% confidence intervals (in parentheses) are indicated at the nodes of the tree in Mya. All nodes had 100% support in 500 bootstrap replicates.



Figure 3: An identity plot of the MHC regions of the Mikado pheasant and the chicken. The chicken MHC sequence was downloaded from GenBank (AB268588). Its nucleotide sequence from 17,978 to 241,251 was aligned against the Mikado pheasant MHC sequence from 2,615 to 229,500 in scaffold208. The gene structure boxes on the horizontal and vertical axes, respectively, represent the gene loci in the Mikado pheasant and the chicken. Boxes with different sizes exhibit different gene locus sizes, and red/blue coloring indicates genes in forward/reverse orientation. The red dots (or lines) on the diagonal indicate that the sequences were aligned in the same orientation, whereas the blue dots indicate alignments with reverse complements. The green dotted lines highlight the sequence of the inverted TAPBP locus and TAP1-TAP2 block. The orange peaks show the read counts on a natural log scale of the gene expression based on our RNA-seq data. The box plot colored in purple indicates d_N/d_S ratios of genes.

 $\ensuremath{\textbf{Table 1: DNA}}$ contigs and scaffolds from the genomic data of the Mikado pheasant

	Contigs	Scaffolds		
Total length	1,054,607,905	1,035,947,982		
Maximum length	195,342	50,275,205		
Number of Ns	0	19,577,473		
Average length	5,050	110,714		
N50*	13,461	11,461,115		
N75*	6,528	5,708,287		
$L50^{\dagger}$	22,195	28		
L75 [†]	50,081	59		
Counts ≥300 bp	208,810	-		
Counts ≥1 kb	123,006	9,357		
Counts ≥5 kb	61,237	1,489		
Counts \geq 10 kb	32,868	928		

 $^{*}\mbox{The N50/N75}$ length is defined as the shortest sequence length at 50%/75% of the genome.

 $^\dagger The$ L50/L75 count is defined as the smallest number of contigs (or scaffolds) that those length sum produces N50/N75.

Values of the genome assembly were calculated using the contigs ${\geq}300$ bp and scaffolds ${\geq}1000$ bp.

tein database for Aves and Reptilians. The statistics of annotated genes in the Mikado pheasant averaged 19.9 kb per gene, 1,625 bp per coding DNA sequence, 164.1 bp per exon, and 2,053 bp per intron (Additional file 1: Table S3), which are similar composition of gene elements in length to other avian species [19]. Of the 15,972 NR annotated proteins, 14,124 proteins were well annotated to the Pfam domains. A total of 5,626 Pfam domains were identified based on our predictions.

Assessment of the assembly quality

The overall DNA mapping rate of the paired-end libraries was >90% for the concordant paired read alignment and >96% for both paired and single read alignment (Additional file 1: Table S4). Thus, the assembly utilized most of the DNA reads. We further examined the per-base alignment coverage. The results (Additional file 1: Fig. S6) showed that most of the genome positions had a coverage between approximately 57- and 121-fold and an average sequence coverage of 88-fold, which is very similar to the sequencing depth of DNA paired-end libraries (98.7x). Thus, our draft genome is well assembled.

To evaluate the quality of the assembled genome [20], the RNA reads were mapped onto the draft genome. The overall alignment rate of both RNA libraries showed that approximately >93% of the reads could be aligned to the scaffolds, indicating that most of the expressed protein-coding genes could be found in the draft genome (Additional file 1: Table S5). Moreover, the Benchmarking Universal Single-Copy Orthologs (BUSCO [BUSCO, RRID:SCR_015008]) [21] benchmark was used to evaluate the genes predicted from the genome assembly (Additional file 1: Table S6). Of the 3,023 single-copy orthologs in the vertebrate lineage, approximately 88.6% of the orthologs were found in our assembly, which is similar to the results obtained in duck (88.6%), turkey (87.5%), and zebra finch (88.8%). These results suggest that a potentially large number of genes, along with their complete structure, could be predicted from the genome.

Genome comparison

To understand the similarities between the Mikado pheasant and the chicken at the genomic level, assembled scaffolds that were longer than 0.25% of the aligned chicken chromosome were selected and plotted onto a syntenic map with an alignment length of at least 3 kb using MUMmer [22]. The identities of each chicken chromosome with the scaffolds of Mikado pheasant were between 86.24% and 89.98%, and the overall coverage was 85.28% (i.e., 855.35 Mb of the assembled scaffolds could be mapped onto the chicken genome; Additional file 1: Table S7). The syntenic relationships between the Mikado pheasant scaffolds and the chicken chromosomes were highly conserved, but a few of the chromosomes could be only partially aligned. In particular, three well-assembled scaffolds, i.e., scaffold14, scaffold69, and scaffold46, were mapped to nearly the full length of chicken chromosomes 15, 23, and 24, respectively. Notably, compared to the scaffolds of the Mikado pheasant, the chicken chromosomes, including chromosomes 6, 11, 18, and 21, were properly aligned but with obvious inversions (Fig. 1A). More stringent conditions were then considered in order to evaluate the alignment of certain scaffolds to multiple chromosomes (e.g., scaffold1 and scaffold45; Fig. 1B); however, further confirmation is required to determine whether this finding represents the actual presence of chromosomal translocations in the Mikado pheasant genome. Additionally, the alignment between the Mikado pheasant scaffolds and the turkey chromosomes provided similar results (Additional file 1: Fig. S7A), but the Mikado pheasant scaffolds were poorly aligned with the zebra finch chromosomes (Additional file 1: Fig. S7B). In general, the Mikado scaffolds displayed high conservation with the genomes of chicken and turkey. We also observed several intrachromosomal inversions and chromosomal translocations. This is the first genomewide analysis to identify multiple intrachromosomal inversions between the Mikado pheasant and chicken genomes.

Phylogenetic relationships of the Mikado pheasant

To compare the protein sequences of the Mikado pheasant against homologous protein families of other birds and organisms, OrthoMCL (OrthoMCL DB: Ortholog Groups of Protein Sequences, RRID:SCR_007839) [23] was used to define the gene families in 10 species. Proteins with sequences that were similar to those of the Mikado pheasant-five birds (i.e., chicken, duck, flycatcher, turkey, and zebra finch), two reptiles (anole lizard and Chinese softshell turtle), and two mammals (human and mouse)-were classified into each gene family. In this way, we obtained 18,220 gene families from 10 species. Next, 5,287 single-copy orthologs that were common across these species were used to construct a Bayesian maximum clade credibility phylogenetic tree and to estimate the time of divergence [24] (Fig. 2). The estimated time of the Mikado pheasant-turkey divergence was 21.4 million years ago (Mya); the divergence time between chicken and the sister clade of the Mikado pheasantturkey was estimated at 28.9 Mya. In the Galliformes order, the Mikado pheasant was found to be more closely related to the turkey than to the chicken. The branches of the Galliformes and duck (76.4 Mya), Passeriformes and Galliformes (105.3 Mya), and anole lizard and Aves (266.3 Mya) displayed divergence times that were similar to those reported in the literature [25-27].

Gene family evolution

To assess the changes in the gene family sizes, a likelihood model was used to examine significant expansions and contractions of gene families, particularly in the Mikado pheasant lineage. Expansions or contractions in gene families indicate that total numbers of genes in a gene family are increasing or decreasing, respectively. The results revealed 311 expanded and 15 contracted gene families compared with the common ancestor of the Mikado pheasant and turkey (Fig. 2). In total, 86 gene ontology (GO) categories were significantly enriched (p < 0.05, empirical test) among the 311 expanded genes. Fifty of these GO categories were further classified into eight main categories, including actin cytoskeleton, morphogenesis, catalytic activity, cell differentiation, binding, metabolism, cytoplasm, and organelle organization and biogenesis (Additional file 2: Table S8). In particular, the gene families involved in oxygen and heme binding (GO:0019825 and GO:0020037, respectively), monooxygenase activity (GO:0004497), and energy metabolism (GO:0046034, ATP metabolic process; GO:0005977, glycogen metabolic process) were substantially expanded in the Mikado pheasant. Conversely, 7 of the 25 GO categories in the contracted gene families were significantly enriched in immune system processes and apoptosis (Additional file 1: Table S9). From the Pfam database [28], 8 of the 75 expanded gene families were annotated as olfactory receptors (Additional file 2: Table S10).

Positive selection

To detect the genes that evolved rapidly due to positive selection under the influence of high elevation (Mikado pheasant) as opposed to low elevation (chicken, turkey, duck, and zebra finch), 7,132 single-copy orthologs were analyzed from 9,038 genes common across the five species (Additional file 1: Fig. S8). According to the branch-site model and the likelihood ratio test, the 889 positively selected genes (PSGs) identified in the Mikado pheasant were mainly enriched in functions such as metabolism (GO:0008152), cell (GO:0005623), and binding (GO:0005488) that belong to biological process, cellular component, and molecular function ontology terms, respectively. We further examined the PSGs involved in metabolism, which constituted the largest number of PSGs and GO functions (Additional file 1: Fig. S9). The 45 PSGs enriched in metabolismrelated functions (p-values < 0.05) were classified according to the GOSlim categories into lipid metabolism (GO:0006629), carbohydrate metabolic processes (GO:0005975), and generation of precursor metabolites and energy (GO:0006091), which included 13, 3, and 2 GO functions, respectively (Additional file 2: Table S11). Of these metabolism-related PSGs, four genes were found to be involved in the inositol phosphate metabolism (map00562; p-value < 0.01) and phosphatidylinositol signaling system (map04070; p-value < 0.05) through a functional enrichment analysis from the Kyoto Encyclopedia of Genes and Genomes (KEGG [KEGG, RRID:SCR_012773]) database (Additional file 1: Table S12).

In addition to metabolism, other high-altitude adaptations were observed, such as response to radiation (GO:0010212, response to ionizing radiation; GO:0010332, response to gamma radiation; GO:0034644, cellular response to ultraviolet radiation; and GO:0071480, cellular response to gamma radiation), DNA repair (GO:0000731, DNA synthesis involved in DNA repair; GO:0045739, positive regulation of DNA repair; and GO:0006284, base-excision repair), and oxygen transport (GO:0016706, oxidoreductase activity; GO:0072593, reactive oxy-

gen species metabolic process; GO:0019825, oxygen binding; and GO:2000377, regulation of reactive oxygen species metabolic process; Additional file 2: Table S13). Moreover, 43 PSGs in the Mikado pheasant were significantly enriched in the categories of lymphocyte activation (GO:0046649; including eight GO terms) and cytokine production (GO:001816; including eight GO terms; Additional file 2: Table S14). We also identified the janus kinase/signal transducer and activator of transcription (Jak-STAT) signaling pathway (map04630; p-value < 0.05), which was enriched in five PSGs (i.e., BCL2, CCND3, IL12RB2, IL23R, and IL7), in the KEGG analysis (Additional file 1: Table S15).

Identification of the MHC-B region of the Mikado pheasant

The MHC is a cluster of genes that are associated with functions such as infectious disease resistance and immune responses in all jawed vertebrates [29]. The MHC B-locus (MHC-B) performs the main MHC functions in the chicken [30, 31]. Based on the above analysis, an assembled scaffold (scaffold208) was almost able to cover the known chicken sequence of the MHC-B contiguous region published by Shiina et al. (GenBank accession: AB268588.1) [32] (Fig. 3). To understand the evolution of the MHC-B genes between the Mikado pheasant and the chicken, the predicted gene loci were manually curated by incorporating evidence from the aligned RNA-sequencing (RNA-seq) data and homologous genes from chicken and turkey using Web Apollo software [33]. After the curation, 39 putative MHC genes of the Mikado pheasant were identified within a 227 kb sequence (Table 2) including seven MHC class II loci (BLB1, TAPBP, BLB2, BRD2, DMA, DMB1, and DMB2), four MHC class I loci (BF1, TAP1, TAP2, and BF2), and five MHC class III loci (C4, CenpA, CYP21, TNXB, and LTB4R1).

Gene loci involved in immunity have been shown to have a higher ratio of nonsynonymous (d_N) to synonymous (d_S) amino acid substitutions due to interactions with rapidly evolving pathogens under selective pressures [34-36]. KIFC1, BTN1, Blec2, BLB1, BLB2, and BF2 had comparatively high d_N/d_S ratios between the Mikado pheasant and the chicken (Table 2). Conversely, the genes with comparatively lower d_N/d_S ratios included TRIM7.2, TRIM41, BRD2, and CenpA. As shown in Fig. 3, the Mikado pheasant and the chicken displayed similarity in the MHC-B region and shared an almost perfect syntenic gene order. Notably, no BLB2 genes were predicted between the TAPBP and BRD2 intergenic regions in the Mikado pheasant MHC-B locus; however, these regions could be detected among the transcripts of our RNA-seq data. A likely explanation for the absence of a prediction of the BLB2-like gene might be the existence of two unsequenced gap regions with a size of 1,098 bp within the TAPBP-BRD2 block (5,931 bp). Since BLB2 is only 792 bp in length, it could reside within the missing sequence. Based on the RNA-seq results, 2.54 million reads were mapped onto 38 MHC-B genes (except for BLB2) of the Mikado pheasant, 27 of which had at least a 1-fold average coverage per nucleotide. Furthermore, 15 genes possessed more than 100-fold average coverage per nucleotide, providing concrete evidence of a reliable prediction. Intriguingly, two gene loci, i.e., TAPBP and the TAP1-TAP2 block, were inversely oriented compared to the chicken sequence.

Evolutionary history of Syrmaticus pheasants

The mitochondrial genome of the Mikado pheasant was assembled based on the short-read libraries. The circular complete genome had a total length of 16,680 bp, including 13 protein-



Figure 4: A phylogenetic tree of *Syrmaticus* pheasants. The divergence times and associated 95% confidence intervals shown in parentheses are given at the branch nodes of the tree in million years ago.

coding genes, 2 rRNAs, 22 tRNAs, and a control region (Additional file 1: Table S16). The average nucleotide composition was 30.52% A, 31.20% C, 13.44% G, and 24.84% T. To investigate the evolutionary history of the genus *Syrmaticus*, which includes five long-tailed pheasants, the phylogeny was reconstructed, and the divergence times were estimated using the mitochondrial genomes. According to molecular clock analysis, the genetic divergence of the Mikado pheasant began approximately 3.47 (2.78–4.71) Mya (Fig. 4). The tree topology is consistent with previous studies [12, 37], and the divergence time suggests that the Mikado pheasant might have originated in the late Pliocene.

Amino acid substitution analysis in Mikado pheasant hemoglobin genes

Living at high elevations directly incurs the challenge of low oxygen availability. Additionally, exposure to low-pressure environments causes oxygen saturation in the arterial blood, thus decreasing and restricting oxygen supplementation to tissues [38]. Certain birds show an increased combined affinity between blood and oxygen via amino acid substitutions in the major hemoglobin [39-41]. To investigate their role in adaptation to high-altitude environments, amino acid substitutions were examined in the Mikado pheasant hemoglobin sequences. By comparing six avian species, an amino acid substitution with different consensus residues was found in the Mikado pheasant (Additional file 1: Fig. S10), and the substitution of alanine with threonine occurred at residue 78 of the alpha-A subunit—the major component of hemoglobin isoforms. The Andean goose, a kind of waterfowl that lives at more than 3,000 meters in the Andes, has been reported to carry the identical substitution [42].

Discussion

In this study, experimental data and statistical approaches were used to evaluate the genome assembly of the Mikado pheasant. Notably, the genome sequence of this species was previously unknown, and this study provides a comparative analysis of various genomes using a large number of tools at different stages for the assembly of the Mikado pheasant genome. While conducting the genome assembly, we used not only MaSuRCA but also assembly tools such as ALLPATHS-LG [43], JR [44], Newbler [45], SGA [46], and SOAPdenovo [47]. All of these assembly tools produced similar draft genome sizes, and MaSuRCA and SGA also showed similar results in terms of the N50 value and the scaffold number (Additional file 1: Table S17). To facilitate the downstream analysis, we used several methods to compare these assembly sets. However, no single assembly tool outperformed the others in terms of the number of annotations for the predicted genes, the quality of the genome compared to that of other birds, and the BUSCO benchmark. In this study, the draft genome assembled using MaSuRCA was selected because it generated dramatically longer scaffolds that displayed a decent score on the BUSCO benchmark and produced proper annotations for the predicted genes. Although scaffolds of the draft genome displayed some degree of fragmentation (Fig. 1A) and showed translocation (Fig. 1B) in certain chicken chromosomes, our approach still provides a practical strategy for wholegenome assembly using only short-read sequencing technology. We assert that the high coverage of our sequencing data, differing library insert sizes, and the use of a combination of tools, such as MaSuRCA and SSPACE for assembly and scaffolding, respectively, contributed to high-quality de novo assembly of the Mikado pheasant genome with a genome length of approximately 1 Gb.

Recent studies have reported phylogenetic tree topologies for the Mikado pheasant and other Galliformes birds [37, 48, 49]; however, these studies relied on small amounts of genomic DNA as supporting evidence. To obtain a highly accurate phylogenetic inference, long DNA sequences are necessary for the reconstruction of a high-resolution tree [50-52]. This study used whole-exome information, with 5,287 single-copy orthologs totaling approximately 8 Mb of coding sequence, to reconstruct the phylogeny and estimate the divergence time among the Mikado pheasant and other birds (Fig. 2). Our results strongly suggest that the Mikado pheasant is more similar to the turkey than the chicken in the Galliformes clade, which is consistent with previous studies [37, 48, 49].

We additionally implemented a comprehensive phylogenetic analysis strategy to obtain information regarding the adaptive mechanisms of the Mikado pheasant to high elevations. Compared to birds that live at low altitudes, both the positive gene selection and gene expansion analyses showed a significant enrichment of genes relevant to energy metabolism (Additional file 2: Tables S8 and S11). This finding was consistent with the prior study that identified similar genes in other species that inhabit the highlands [53]. Moreover, the four metabolism-related PSGs (i.e., INPP5A, INPP5J, PI4KB, and PLCE1) that were involved in the inositol phosphate metabolism and phosphatidylinositol signaling system (Additional file 1: Table S12) were previously reported to be enriched in Tibetan pigs living at high altitudes [54]. Of these genes, INPP5A and INPP5J play a role in the hydrolysis of inositol polyphosphates [55], PI4KB is a phosphatidylinositol kinase that induces phosphorylation reactions [56], and PLCE1, which is a phospholipase enzyme, regulates gene expression, cell growth, and differentiation [57]. Another robust signal of its adaptation to high altitude was obtained from genes significantly associated with expansion of and positive selection for the enhancement of hemoglobin binding and oxygen transport (Additional file 2: Tables S8 and S13). Furthermore, for both the Mikado pheasant and Andean goose, an amino acid substitution was identified in the hemoglobin alpha-A subunit (Additional file 1: Fig. S10). The substitution of threonine at this position has recently been shown to cause an increase in the molecular volume, which might enhance the solubility of hemoglobin and facilitate adaptation to desiccating and high-altitude environments [42]. Through gene expansion, the genes of the Mikado pheasant that are involved in skeletal and cardiac muscle fiber development (Additional file 2: Table S8) and the enhanced functions of the additional GO terms implied that the biomass of the Mikado pheasant could be effectively produced in mountain-

Table 2: Coding sequences of MHC-B genes in the Mikado pheasant and comparisons with the chicken

Mikado pheasant				Chicken							
Gene	Position	Strand	Gene length	Amino acid length	Exon	Aligned base	Nucleotide identity (%)	Aligned amino acid	Amino acid identity (%)	Amino acid substitu- tions	d_N/d_S^*
KIFC1	2615-5304	+	1,140	380	7	1,131	91.76	377	90.53	33	0.8669
Blec3	8997-11221	-	552	183	5	507	82.43	168	78.14	25	0.3821
Bzfp3	12 126-18 213	+	1,449	482	13	1,569	85.8	522	83.62	40	0.1884
TRIM7.2	19 507–24 562	-	1,518	505	7	1,518	95.98	505	98.61	7	0.0391
Bzfp2†	27 027–29 946	+	1,368	455	4	1,396	70.41	N/A	N/A	N/A	0.2438
Bzfp1	31 049–33 298	-	1,425	474	2	1,426	88.23	471	86.79	54	0.1900
44G24.1	37 266–37 673	-	408	136	1	408	85.78	136	80.15	27	0.2762
IL4I1	42730-46759	+	1,578	525	6	1,572	92.25	523	93.75	25	0.1011
TRIM7.1	51 325–62 131	-	1,758	585	8	1,767	92.49	588	92.69	40	0.1545
HEP21	63 362–64 247	-	324	107	3	324	93.52	107	91.59	9	0.2148
TRIM39.2	70 980–74 640	-	1,392	464	6	1,389	93.68	463	94.61	24	0.1167
TRIM27.2	76 988–80 522	+	1,431	476	7	1,431	94.13	476	92.23	37	0.2415
TRIM39.1	81 560-85 449	-	798	266	5	798	93.23	266	91.35	23	0.2753
TRIM27.1	86 518–90 228	-	1,485	495	7	1,485	94.48	495	94.34	28	0.1715
TRIM41	91 918–96 605	+	1,656	551	7	1,770	89.58	589	91.71	7	0.0375
GNB2L1	98 038–101 512	-	954	317	8	954	96.86	317	100	0	N/A
BTN1	103 411-114 264	+	930	309	8	939	74.64	339	57.26	96	0.8357
BTN2	117 466–120 157	+	1,461	487	7	1,481	90.41	469	83.37	52	0.3996
BG1	124 105–125 436	-	549	183	3	546	91.99	182	87.98	21	0.5591
Blec2	131 358–133 021	-	579	192	5	579	86.32	190	71.88	52	0.9375
Blec1	135 818–137 846	+	567	188	5	567	92.59	188	88.3	22	0.4683
BLB1	138 411–139 729	-	339	112	3	345	83.38	113	42.98	62	0.7904
TAPBP	140 657–144 216	+	1,293	430	8	1,293	92.19	430	89.77	44	0.3179
BLB2 [‡]	N/A	N/A	792	263	N/A	792	92.93	263	85.93	37	1.4489
BRD2	150 146–156 295	-	2,976	991	13	3,078	86.85	776	75.28	30	0.0306
DMA	160 545–162 778	+	789	263	4	789	92.65	263	89.73	27	0.4528
DMB1	163 010–165 184	+	930	310	6	930	91.29	310	86.45	42	0.4978
DMB2	165 617–168 363	+	768	256	5	768	92.71	256	92.58	19	0.1622
BF1	169 254–170 740	+	996	331	5	1,001	83.66	345	64.12	95	0.7116
TAP2	172 793–176 021	-	2,100	700	9	2,100	92.48	700	93.14	48	0.1675
TAP1	176 574–180 981	+	1,752	584	11	1,739	93.21	580	92.81	38	0.2191
BF2	181 900–184 038	-	1,530	509	6	1,213	62.28	326	57.39	119	0.8157
C4	185 102–199 258	+	5,031	1,676	40	4,998	93.33	1,665	93.2	101	0.1974
CenpA	199 593–200 795	+	396	131	4	396	96.72	131	99.24	1	0.0324
CYP21	201 291–205 141	+	1,431	477	11	1,431	92.67	477	94.13	28	0.7109
TNXB	209 524–215 604	-	2,472	824	10	2,496	92.14	832	92.34	50	0.2002
LTB4R1	221 450-222 538	+	1,089	363	1	1,089	94.12	363	94.49	20	0.1954
CD1A2	223 740-225 788	-	1,044	348	6	1,044	92.24	348	87.93	42	0.3796
CD1A1	227 030-229 500	-	1,122	374	6	1,122	93.4	374	90.64	35	0.3294

KIFC1, kinesin family member C1; Blec, C-type lectin-like receptor; Bzfp, B-locus zinc finger-like protein; TRIM, tripartite motif containing protein; 44G24.1, histone H2B-like protein; IL4I1, interleukin 4 induced 1; HEP21, hen egg protein 21 kDa; GNB2L1, guanine nucleotide binding-like protein; BTN, B-butyrophilin protein; BG1, BG-like antigen; CD1A1/A2, CD1-like proteins.

 $^{*}d_{N}/d_{S} =$ ratio of nonsynonymous (d_N) to synonymous (d_S) substitutions.

[†]Defined as a pseudogene in chicken.

*No predicted result was identified from the DNA assembly. The transcript sequence was alternatively derived from the transcriptome assembly by RNA-seq.

ous regions without nourishment, hence strongly suggesting the existence of an adaptive mechanism for high altitudes [58]. Finally, the PSGs in the radiation response, immune response, and DNA repair categories (Additional file 2: Tables S13 and S14) may reflect the increased resistance of the Mikado pheasant to long-term ultraviolet radiation exposure through the induction of cytokine production [59] and lymphocyte activation [60] and DNA repair processes. Some of these PSGs were also involved in the Jak-STAT signaling pathway (Additional file 1: Table S15), which participates in chemical signal transmission and induces cellular stress responses, such as immunity, apoptosis [61, 62], and hypoxia [63]. All of these results provide wider support for

the adaptive evolution of the Mikado pheasant. To sum up, this study reveals the high-altitude adaptation mechanisms of the Mikado pheasant at the genomic level. However, there are some adaptive mechanisms for high altitude that happen via changes in regulatory regions that modulate the levels of gene expression [64-66]. We believe that this is an intriguing topic and worthy of further research to be undertaken in the future.

In this work, we annotated and curated the MHC-B gene loci in the Mikado pheasant, which is important for assessing the adaptive mechanisms associated with endangered species, because variations in gene number in the MHC cluster could be caused by exposure to pathogens or diseases [67, 68]. The genome of the Mikado pheasant contains a number of MHC-B genes, and inversions were observed in the TAPBP locus and the TAP1-TAP2 block (Fig. 3) compared to the chicken genome; an inverse orientation of the TAP1-TAP2 block was also detected compared to the turkey genome (Additional file 1: Fig. S11). A similar conversion at the MHC locus in Galliformes has been reported in previous studies [29, 34, 69]. We further observed a Blec2-like sequence with an inverse orientation located within the BG1-Blec2 region in the Mikado pheasant. We inferred that this region is likely similar to the Blec4 pseudogene of the chicken and highly similar to Blec2 [32].

In this study, we not only sequenced the whole genome of a bird of the Syrmaticus genus but also completed the full mitochondrial genome. Before whole-genome sequences were available, mitochondrial sequences were widely utilized in molecular phylogenetic analyses of the genus of Gallus [70, 71]. Based on the assembly of the Mikado pheasant and the other four available sequences, we reconstructed a phylogenetic tree and provide a completely sequenced mitochondrial genome for five long-tailed pheasants. The topology of our reconstructed tree (Fig. 4) is consistent with results from a previous study [12]. However, the time of divergence was estimated to be earlier than the previously reported time [12] for the Mikado pheasant, which might have been due to the use of a few mitochondrial or nuclear genes rather than the complete mitochondrial genome. The reconstructed tree showed a potential migration pathway of these pheasants. The ancestors of the Mikado pheasant, which have been described to have migrated to the island of Taiwan, separated from the lineage of the copper pheasant (Syrmaticus soemmerringii ijimae). The copper pheasant is a pheasant indigenous to Japan, whose ancestors might have separated from the lineage of the Reeves's pheasant (Syrmaticus reevesii) that has inhabited northern China. The ancestors of Elliot's pheasant (Syrmaticus ellioti) and Mrs. Hume's pheasant (Syrmaticus humiae) have branched from the Mikado pheasant, then separated into two present kinds of pheasants that have alternatively roosted in the mountainous forests of southeastern and southwestern China, respectively. According to paleogeographical reports, Taiwan was formed approximately 4-5 Mya and attained its modern topography approximately 3 Mya [72]. The sea level was lower during the glacial periods, and Taiwan might have been connected to the mainland [73]. Our results suggest that the evolutionary history of the Mikado pheasant might have included ancestors that migrated from the north toward Taiwan approximately 3.47 Mya and consequently were isolated by the Taiwan Strait during the warm interglacial periods during the early Pleistocene.

Currently, there is no nuclear genome data available for the copper pheasant, so unfortunately, incorporating all five longtailed pheasants into our analysis using nuclear genomes is impossible at present. For the other four pheasants, however, Wang et al. [37] used six nuclear intron and two mitochondrial gene sequences to construct a phylogenetic tree, and its topology was consistent with our result. Our estimate of the divergence time was more precise, considering that we employed complete mitochondrial genomes in the reconstruction of a high-resolution tree for the Syrmaticus genus instead of a few mitochondrial genes. Our estimated divergence time is also supported by the paleogeographical report of Taiwan island formation. Despite these corroborations of the proposed tree topology and estimated divergence time, the use of only mitochondrial data may be considered as a potential limitation. Going forward, it will be necessary to analyze the nuclear genome to obtain further insights into the evolution history of long-tailed pheasants.

Materials and Methods

Sample preparation and sequencing

Blood samples were collected from a single female Mikado pheasant living in central Taiwan; then, genomic DNA was extracted, and two paired-end libraries (280 bp and 480 bp; average read length, 151 bp) and five mate pair libraries (1, 3, 5, 7, and 10 kb; average read length, 101 bp) were constructed according to the manufacturer's protocol. In addition, two RNA-seq libraries from two male Mikado pheasants' blood samples were prepared for the purpose of draft genome assessment and gene prediction (Additional file 1: Table S1). The DNA libraries were sequenced using the HiSeq platform (Illumina Inc., San Diego, CA, USA), and the RNA libraries were sequenced using the HiScanSQ and HiSeq platforms.

De novo genome assembly

The quality of the raw reads was examined using FastQC (FastQC, RRID:SCR_014583), version 0.10.1. Trimmomatic (Trimmomatic, RRID:SCR_011848), version 0.30 (parameters: "ILLUMINACLIP: TruSeq3-PE.fa:2:30:15 SLIDINGWINDOW:4:20 MINLEN:100") [74] and NextClip (version 1.3.1) [75] with default parameters were used to trim sequencing reads. Genome assembly into contigs was performed using MaSuRCA (version 2.3.2) [15] with settings based on the instruction manual. ALLPATHS-LG (ALLPATHS-LG, RRID:SCR_010742, version 49722) [43], Newbler (version 2.9) [45] both with default parameters, JR (version 1.0.4; parameters: "-minOverlap 60 -maxOverlap 90 -ratio 0.3") [44], SGA (version 0.10.13; parameters: "assemble -m 125 -d 0.4 -g 0.1 -r 10 -l 200") [46], and SOAPdenovo (version 2.04; parameters: "-K 47 -R") [47] were also used to assemble contigs. We employed SSPACE (SSPACE, RRID:SCR_005056, version 3.0; parameter: "-z 300") [76] to construct scaffolds for the draft genome. In this step, mate pair libraries with 35 bases from the 5' end of both reads were used for scaffolding. Scaffold sequences shorter than 300 bp were then excluded from the final assembly. The statistical results of the assembly were estimated using QUAST (version 3.2) [77].

To examine sequencing reads for potential contamination, we used Kraken (version 1.0) [78] with the standard Kraken database to check the paired-end DNA libraries. Classified reads reported by Kraken were further examined using our proposed pipeline (Additional file 1: Fig. S12). Briefly, we employed Bowtie 2 (Bowtie, RRID:SCR_005476; version 2.3.0) [79] to align these classified reads against the chicken genome reference (Galgal 5.0) downloaded from Ensembl (release 90), collecting unmapped reads and using Bowtie 2 again to align them against the assembled genome of the Mikado pheasant. We then took those reads mapped onto the Mikado pheasant genome and performed Basic Local Alignment Search Tool N (BLASTN) alignment against the nonredundant nucleotide sequences database, downloaded from NCBI's FTP site (on Nov. 16, 2017), using parameters "outfmt "6 std staxids" -max_target_seqs 1 -evalue 1E-10." Next, we collected reads with alignment length \geq 100 bp (i.e., two thirds of read length), filtering out the reads that matched an avian species or with a read count <50 in a species. The remaining reads were counted and the contaminated scaffolds calculated by applying a cutoff of a read count >20 on a given scaffold. Finally, we removed 31 contaminated scaffolds with 12,587 bp (\sim 0.001% of the total length) from the assembled genome.

Evaluation of assembly quality

Several metrics were used to evaluate the assembly quality, including the number and length distribution of the scaffold sequences, the mapping rate of the paired-end DNA reads and RNA reads, the per-base coverage of the DNA read mapping, and the coverage of universal single-copy orthologs provided by BUSCO (version 1.21). To evaluate the mapping rate of the reads and perbase coverage, the paired-end DNA reads and RNA reads were aligned against the assembled scaffolds using Bowtie 2 (version 2.2.4) and STAR [80], respectively. Briefly, scaffolds were mainly assembled from the paired-end DNA reads; the higher mapping rate of the paired-end DNA reads suggests a higher degree of the final assembly covering the raw reads. Taking the RNA sequencing reads from two individual Mikado pheasants and observing the mapping rate is another approach for assessing the completeness of the assembly. The per-base DNA read coverage was calculated using BEDTools (BEDTools, RRID:SCR_006646), version 2.23.0 [81]. For each base, the expected coverage should be close to the sequencing depth of the paired-end reads (approximately 98.7x). The BUSCO benchmark is a single-copy ortholog set derived from the species of a major lineage. The gene models predicted from the draft genome in the Mikado pheasant were compared with the lineage of vertebrates (3,023 single-copy orthologs in total) provided by BUSCO. Protein sequences from the chicken, duck, turkey, and zebra finch were also evaluated for comparison.

Genome comparison

To compare the genome of the Mikado pheasant with that of other avian species, we retrieved the whole-genome sequences of the chicken (Galgal4), turkey (UMD2), and zebra finch (taeGut3.2.4) from the Ensembl database. Using the genome-wide sequence aligner MUMmer (version 3.23), the chromosome-level differences and similarities among the species were investigated and visualized. The structural variants among the species were further reported using the "show-diff" utility in MUMmer. The chord diagrams of the alignment were generated using Circos [82].

Gene prediction and annotation

First, RepeatMasker (version 4.0.5; parameter: "-species chicken"), including rmblastn (version 2.2.23+) as the search engine, RepBase (version 20140131), and RM database (version 20140131), were applied to screen the scaffolds for interspersed repeats and low-complexity regions in the DNA sequences, and the masked genome was used for further gene prediction. Then, homology-based, RNA-seq, and ab initio prediction approaches were used to identify protein-coding genes and build a consensus gene set that included all predicted genes. For the homology protein sequence alignment, the protein sequences of the chicken (Galgal4), turkey (UMD2), duck (BGI_duck_1.0), and zebra finch (taeGut3.2.4) were collected from Ensembl. The protein sequence alignments were performed using Exonerate (version 2.2.0) [83]. All RNA-seq reads were aligned against the repeat-masked genome using TopHat2 [84], which generated evidence of splice sites, introns, and exons. Additionally, Trinity (Trinity, RRID:SCR_013048), version 2.0.6, [85], was utilized to assemble transcripts, and PASA (version 2.0.0) [86] was used to group alternatively spliced isoforms. For the ab initio gene prediction, the standard Augustus (version 3.0.3) pipeline was used to yield potentially predicted genes with evidence from both homologous proteins and RNA-seq. Next, the consensus gene set was determined by consolidating the three types of gene prediction using EVidenceModeler (version 1.1.1). Finally, the gene annotations were defined based on the best sequence alignment against NCBI NR proteins in Aves and Reptilians using BLASTP (version 2.2.29+), with the following criteria: identity \geq 30%, alignment length \geq 80 bp, and E-value \leq 1e–5. For the protein domain identification, we annotated the domains using HMMER (version 3.1b2) [87] by scanning the Pfam database (version 30.0).

For MHC-B annotation and curation, we first took the scaffold208 sequence and used MAKER (version 2.31.8) [88] to predict the potential gene structures of MHC-B genes. Next, the RNA-seq libraries from the Mikado pheasant and the homologous protein sequences from chicken and turkey were aligned to these predicted regions. Finally, we used Web Apollo (version 2.0.3), a web-based and visualization tool for curation and annotation, to manually curate these genes according to the alignment evidence.

Gene families

To identify gene families, the protein-coding genes of five birds (i.e., Gallus gallus, Meleagris gallopavo, Anas platyrhynchos, Taeniopygia guttata, and Ficedula albicollis) and four additional species (Anolis carolinensis, Pelodiscus sinensis, Homo sapiens, and Mus musculus) were downloaded from Ensembl (release 82). The sequence of the longest isoform was selected to represent the gene for each species, despite the presence of protein isoforms. The all-vs-all BLASTP was applied to align all protein sequences (including those of the Mikado pheasant) of the 10 species and 5 birds (excluding flycatcher) with E-value thresholds less than 1e-5 and 1e-20, respectively. Then, 18,220 gene families (including 5,287 single-copy orthologs) were obtained from the 10 species, and 13,436 gene families (including 7,132 single-copy orthologs) were obtained from the 5 birds by OrthoMCL (version 2.0.9) using default parameters. In the analysis of the 10 species, 15,161 genes of the Mikado pheasant were grouped into 12,549 gene families. In the analysis of the five avian species, 14,375 Mikado pheasant genes were grouped into 12,078 gene families. Next, MUSCLE (MUSCLE, RRID:SCR_011812), version 3.8.1551 [89], was used with default parameters for the multiple sequence alignment of the converted coding DNA sequences from singlecopy orthologs, and Gblocks (version 0.91b; parameters: "-t = d -b4 = 5 - b5 = h - e = cln") [90] was used to remove the poorly aligned regions. After trimming, the genes from each species were concatenated using the same order to reconstruct the phylogenies and evaluate the divergence time. The concatenated sequences were used to build a phylogenetic tree using RAxML (RAxML, RRID:SCR_006086), version 8.2.4 [91], via a maximum likelihood search with 500 bootstrap replicates; then, the divergence time was analyzed using BEAST (BEAST, RRID:SCR_0 10228), version 2.3.2, with the GTR+I+ Γ model, which is the best substitution model selected by Modeltest (version 3.7) and PAUP* (version 4.0a150) [92]. Four nodes were chosen as the fossil calibration points from the TimeTree database [93], including human-chicken (311.9 Mya), anole lizard-chicken (279.7 Mya), Chinese softshell turtle-chicken (253.7 Mya), and human-mouse (89.8 Mya). The phylogenetic tree was generated using the Strap R package [94]. To identify the gene families with an expansion or contraction between the Mikado pheasant and other species, CAFE (version 3.1) [95] was used to estimate the rates of gene family evolution from the observed gene numbers in each family and the given phylogenetic tree. A p-value <0.05 was used to indicate significant changes in the gene family size.

Examination of genes under positive selection and enrichment analysis

To determine the genes that underwent positive natural selection in the Mikado pheasant, CODEML from PAML (PAML, RRID:SCR_014932), version 4.8 [96], was applied to the branchsite model to investigate the genes in positively selected sites of the Mikado pheasant. For the branch-site model, we implemented likelihood ratio tests to determine the statistical significance of positive selection for testing a null model (model = 2, NSsites = 2, fix.omega = 1, and omega = 1) against an alternative model (model = 2, NSsites = 2, and fix.omega = 0). Consequently, the false discovery rates (FDRs) were computed with a cutoff of 0.05 to adjust for multiple testing using the Benjamini-Hochberg procedure.

The GO annotations of four birds (i.e., chicken, duck, turkey, and zebra finch) retrieved from the Ensembl BioMart were used to characterize the functions of the identified orthologs. A hypergeometric test was performed to identify significant GO functions in these orthologs. However, the raw p-values of the hypergeometric tests can easily be affected by the number of genes [97]; therefore, to address the underlying bias of the hypergeometric distribution, we further calculated empirical p-values [98]. The empirical p-values were determined through 100K simulated datasets by ranking the hypergeometric probability of enriched functional categories compared with the null baseline probabilities. The null baseline probability was established by randomly selecting a group of genes containing an equal number of PSGs with an FDR <0.05 for the branch-site model. For massively enriched GO terms with similar functions, CateGOrizer [99] was used to classify the genes into basic categories. ClueGO [100] with the hypergeometric test and a Bonferroni adjustment were performed to enrich the KEGG pathways [101].

Mitochondrial genome assembly

Geneious (version 8.1.5) [102] was utilized with the default settings to assemble the whole mitochondrial genome. First, the reads were mapped to the four available Syrmaticus mitochondrial genomes from GenBank (AB164622.1 - AB164625.1). The mapped reads were collected and then used for the further assembly of the mitochondrial genome of the Mikado pheasant. The genes were identified using MITOS [103] and curated by comparison with known sequences of other long-tailed pheasants from GenBank. The phylogenetic reconstruction and estimation of the divergence times among the five long-tailed pheasants were achieved using BEAST with the GTR+G model, which was selected as the best nucleotide substitution model by Modeltest and PAUP*. We added two nodes as the fossil calibration points according to the TimeTree database, including Elliot's pheasant-Reeves's pheasant (11.1 Mya) and Elliot's pheasant-Mrs. Hume's pheasant (0.2 Mya). A calibrated Yule speciation process was implemented in the analysis using BEAST. In the Markov chain Monte Carlo analysis, the chain length utilized 10 million generations.

Availability of supporting data

Data for the S. mikado genome has been deposited in the Gen-Bank/EMBL/DDBJ Bioproject database under the project number PRJNA389983. Raw genomic and transcriptomic sequence datasets were deposited in the Sequence Read Archive under the accession number SRP108966. Other supporting data, including the draft genome, annotations, alignments, phylogenetic trees, and scripts, are available via the *GigaScience* repository, GigaDB [104].

Ethics approval and consent to participate

All experimental procedures and sample collection methods in this study involving Mikado pheasants were conducted according to the Wildlife Conservation Act (amendment on July 8, 2009, Taiwan) and were approved by the Council of Agriculture, Executive Yuan, Taipei, Taiwan with issue 1021700417.

Additional files

Additional file 1: Supplementary Figures S1-S12 and supplementary Tables S1-S7, S9, S12, and S15-S17. Additional file 2: Supplementary Tables S8, S10-S11, and S13-S14.

Abbreviations

BLAST: Basic Local Alignment Search Tool; BUSCO: Benchmarking Universal Single-Copy Orthologs; FDR: false discovery rate; GO: gene ontology; KEGG: Kyoto Encyclopedia of Genes and Genomes; MHC: major histocompatibility complex; Mya: million years ago; NCBI: National Center for Biotechnology Information; NR: nonredundant; PSG: positively selected gene; RNA-seq: RNAsequencing.

Competing interests

The authors declare that they have no competing interests.

Funding

This work was supported by a grant from Taipei Zoo (No. 13, 2015 Animal Adoption Programs of Taipei Zoo). The funder had no role in the design, collection, analysis, or interpretation of the data; writing the manuscript; or the decision to submit the manuscript for publication.

Author Contributions

E.Y.C., C.-Y.C., M.-H.T., S.-T.D., and H.L. conceived the project. E.Y.C., C.-Y.C., M.-H.T., and E.-C.L. managed and coordinated the project. M.-H.T., S.-T.D., and E.-C.L. performed animal work and prepared biological samples. T.-P.L. and L.-C.L. designed bioinformatics and evolutionary analyses. C.-Y.L., P.-H.H., and K.-Y.L. performed genome assembly. P.-H.H. performed assessment of the assembly quality. C.-Y.L. and P.-H.H. performed gene prediction and annotation. C.-Y.L. and P.-H.H. performed evolutionary analysis. C.-Y.L. performed mitochondrial genome assembly and gene annotation and curated the MHC-B gene loci. Y.-F.L. wrote a visualization program for displaying MHC-B genes. C.-Y.L., P.-H.H., and A.C. wrote the manuscript. A.C., M.-H.T., and C.-Y.C. commented on the draft and revised the manuscript; M.-H.T., C.-Y.C., and E.Y.C. cosupervised the study. All authors read and approved the final manuscript.

Acknowledgements

We thank Dr. Mei-Yeh Lu and the High Throughput Genomics Core at the Biodiversity Research Center, Academia Sinica, for the NGS library constructions and Illumina sequencing. We are also grateful to the Computer and Information Networking Center, National Taiwan University, and the National Center for High-Performance Computing for the support of the highperformance computing facilities. We thank Dr. Chih-Ming Hung and Dr. Chen Siang Ng for discussion and comments on the phylogenetic analysis, Dr. Chia-Yang Cheng and Tai-Rong Hong for assistance in data analyses, Dr. Melissa Stauffer and Dr. Yao-Yin Chang for assistance with editing the manuscript, and Yu-Cheng Hsieh for providing photos of the Mikado pheasant.

References

- Bridgman CL. Habitat use, distribution and conservation status of the mikado pheasant (Syrmaticus mikado) in Taiwan, The University of Tennessee, Knoxville; 2002.
- 2. Severinghaus SR. A study of the Swinhoe's and Mikado pheasants in Taiwan with recommendations for their conservation, Cornell University, Ithaca; 1977.
- McGowan PJ, Garson PJ. Pheasants: status survey and conservation action plan 1995–1999, IUCN,Gland, Switzerland; 1995.
- Fuller RA, Garson PJ. Pheasants: status survey and conservation action plan 2000–2004, IUCN,Gland, Switzerland; 2000.
- Yu L, Wang GD, Ruan J, et al. Genomic analysis of snub-nosed monkeys (*Rhinopithecus*) identifies genes and processes related to high-altitude adaptation. Nat Genet 2016;48(8):947–52.
- Le Duc D, Renaud G, Krishnan A, et al. Kiwi genome provides insights into evolution of a nocturnal lifestyle. Genome Biology 2015;16:147.
- 7. Li S, Li B, Cheng C, et al. Genomic signatures of nearextinction and rebirth of the crested ibis and other endangered bird species. Genome Biology 2014;15(12):557.
- Hung CM, Shaner PJ, Zink RM, et al. Drastic population fluctuations explain the rapid extinction of the passenger pigeon. Proc Natl Acad Sci U S A 2014;111(29):10636–41.
- 9. Qiu Q, Zhang G, Ma T, et al. The yak genome and adaptation to life at high altitude. Nat Genet 2012;44(8):946–9.
- Li R, Fan W, Tian G, et al. The sequence and de novo assembly of the giant panda genome. Nature 2010;463(7279):311–7.
- Qu Y, Zhao H, Han N, et al. Ground tit genome reveals avian adaptation to living at high altitudes in the Tibetan plateau. Nat Commun 2013;4:2071.
- 12. Zhan XJ, Zhang ZW. Molecular phylogeny of avian genus *Syrmaticus* based on the mitochondrial cytochrome B gene and control region. Zoolog Sci 2005;**22**(4):427–35.
- 13. Lee P, K Lue, Hsieh J, et al. A wildlife distribution database in Taiwan. Council of Agriculture, Taipei. 1998.
- Chikhi R, Medvedev P. Informed and automated kmer size selection for genome assembly. Bioinformatics 2014;30(1):31–7.
- Zimin AV, Marcais G, Puiu D, et al. The MaSuRCA genome assembler. Bioinformatics 2013;29(21):2669–77.
- 16. Smit AF, Hubley R, Green P. RepeatMasker Open-3.0. 1996.
- Stanke M, Diekhans M, Baertsch R, et al. Using native and syntenically mapped cDNA alignments to improve de novo gene finding. Bioinformatics 2008;24(5):637–44.

- Haas BJ, Salzberg SL, Zhu W, et al. Automated eukaryotic gene structure annotation using EVidenceModeler and the Program to Assemble Spliced Alignments. Genome Biology 2008;9(1):R7.
- Zhang G, Li B, Li C, et al. Comparative genomic data of the Avian Phylogenomics Project. GigaScience 2014;3(1):26.
- Lee C-Y, Chiu Y-C, Wang L-B, et al. Common applications of next-generation sequencing technologies in genomic research. Translational Cancer Research 2013;2(1):33–45.
- Simao FA, Waterhouse RM, Ioannidis P, et al. BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs. Bioinformatics 2015;31(19):3210–2.
- Kurtz S, Phillippy A, Delcher AL, et al. Versatile and open software for comparing large genomes. Genome Biology 2004;5(2):R12.
- Li L, Stoeckert CJ, Jr., Roos DS. OrthoMCL: identification of ortholog groups for eukaryotic genomes. Genome Research 2003;13(9):2178–89.
- Drummond AJ, Suchard MA, Xie D, et al. Bayesian phylogenetics with BEAUti and the BEAST 1.7. Mol Biol Evol 2012;29(8):1969–73.
- Lu L, Chen Y, Wang Z, et al. The goose genome sequence leads to insights into the evolution of waterfowl and susceptibility to fatty liver. Genome Biology 2015;16:89.
- Jiang L, Wang G, Peng R, et al. Phylogenetic and molecular dating analysis of Taiwan Blue Pheasant (Lophura swinhoii). Gene 2014;539(1):21–9.
- Cai Q, Qian X, Lang Y, et al. Genome sequence of ground tit *Pseudopodoces humilis* and its adaptation to high altitude. Genome Biology 2013;**14**(3):R29.
- Finn RD, Mistry J, Tate J, et al. The Pfam protein families database. Nucleic Acids Res 2010;38(Database issue):D211– 22.
- Wang B, Ekblom R, Strand TM, et al. Sequencing of the core MHC region of black grouse (*Tetrao tetrix*) and comparative genomics of the galliform MHC. BMC Genomics 2012;13:553.
- Kaufman J, Milne S, Gobel TW, et al. The chicken B locus is a minimal essential major histocompatibility complex. Nature 1999;401(6756):923–5.
- Kaufman J, Volk H, Wallny HJ. A "minimal essential Mhc" and an "unrecognized Mhc": two extremes in selection for polymorphism. Immunol Rev 1995;143:63–88.
- 32. Shiina T, Briles WE, Goto RM, et al. Extended gene map reveals tripartite motif, C-type lectin, and Ig superfamily type genes within a subregion of the chicken MHC-B affecting infectious disease. J Immunol 2007;178(11):7162–72.
- Lee E, Helt GA, Reese JT, et al. Web Apollo: a web-based genomic annotation editing platform. Genome Biology 2013;14(8):R93.
- Chaves LD, Krueth SB, Reed KM. Defining the turkey MHC: sequence and genes of the B locus. J Immunol 2009;183(10):6530–7.
- Emerman M, Malik HS. Paleovirology–modern consequences of ancient viruses. PLoS Biol 2010;8(2):e1000301.
- Sawyer SL, Wu LI, Emerman M, et al. Positive selection of primate TRIM5alpha identifies a critical species-specific retroviral restriction domain. Proc Natl Acad Sci U S A 2005;102(8):2832–7.
- Wang N, Kimball RT, Braun EL, et al. Assessing phylogenetic relationships among galliformes: a multigene phylogeny with expanded taxon sampling in Phasianidae. PLoS One 2013;8(5):e64312.

- Powell FL, Shams H, Hempleman SC, et al. Breathing in thin air: acclimatization to altitude in ducks. Respir Physiol Neurobiol 2004;144(2–3):225–35.
- Monge C, Leon-Velarde F. Physiological adaptation to high altitude: oxygen transport in mammals and birds. Physiol Rev 1991;71(4):1135–72.
- Weber RE, Jessen TH, Malte H, et al. Mutant hemoglobins (alpha 119-Ala and beta 55-Ser): functions related to high-altitude respiration in geese. J Appl Physiol (1985) 1993;75(6):2646–55.
- Jessen TH, Weber RE, Fermi G, et al. Adaptation of bird hemoglobins to high altitudes: demonstration of molecular mechanism by protein engineering. Proc Natl Acad Sci U S A 1991;88(15):6519–22.
- 42. McCracken KG, Barger CP, Sorenson MD. Phylogenetic and structural analysis of the HbA (alphaA/betaA) and HbD (alphaD/betaA) hemoglobin genes in two high-altitude waterfowl from the Himalayas and the Andes: Bar-headed goose (Anser indicus) and Andean goose (Chloephaga melanoptera). Mol Phylogenet Evol 2010;56(2):649–58.
- Gnerre S, Maccallum I, Przybylski D, et al. High-quality draft assemblies of mammalian genomes from massively parallel sequence data. Proc Natl Acad Sci U S A 2011;108(4):1513–8.
- Chu TC, Lu CH, Liu T, et al. Assembler for de novo assembly of large genomes. Proc Natl Acad Sci U S A 2013;110(36):E3417–24.
- Margulies M, Egholm M, Altman WE, et al. Genome sequencing in microfabricated high-density picolitre reactors. Nature 2005;437(7057):376–80.
- Simpson JT, Durbin R. Efficient de novo assembly of large genomes using compressed data structures. Genome Research 2012;22(3):549–56.
- Luo R, Liu B, Xie Y, et al. SOAPdenovo2: an empirically improved memory-efficient short-read de novo assembler. GigaScience 2012;1(1):18.
- Eo SH, Bininda-Emonds OR, Carroll JP. A phylogenetic supertree of the fowls (Galloanserae, Aves). Zoologica Scripta 2009;38(5):465–81.
- Kimball RT, Mary CM, Braun EL. A macroevolutionary perspective on multiple sexual traits in the phasianidae (galliformes). Int J Evol Biol 2011;2011:423938.
- Dunn CW, Hejnol A, Matus DQ, et al. Broad phylogenomic sampling improves resolution of the animal tree of life. Nature 2008;452(7188):745–9.
- 51. Naylor GJ, Brown WM. Structural biology and phylogenetic estimation. Nature 1997;**388**(6642):527–8.
- Rosenberg MS, Kumar S. Incomplete taxon sampling is not a problem for phylogenetic inference. Proc Natl Acad Sci U S A 2001;98(19):10751–6.
- Li M, Tian S, Jin L, et al. Genomic analyses identify distinct patterns of selection in domesticated pigs and Tibetan wild boars. Nat Genet 2013;45(12):1431–8.
- 54. Zhang B, Qiangba Y, Shang P, et al. A comprehensive MicroRNA expression profile related to hypoxia adaptation in the Tibetan pig. PLoS One 2015;**10**(11):e0143260.
- Hsu F, Mao Y. The structure of phosphoinositide phosphatases: insights into substrate specificity and catalysis. Biochim Biophys Acta 2015;1851(6):698–710.
- Rapoport SI, Primiani CT, Chen CT, et al. Coordinated expression of phosphoinositide metabolic genes during development and aging of human dorsolateral prefrontal cortex. PLoS One 2015;10(7):e0132675.
- 57. Tan J, Yu CY, Wang ZH, et al. Genetic variants in the inositol

phosphate metabolism pathway and risk of different types of cancer. Sci Rep 2015;5:8473.

- 58. Zhu L, Li M, Li X, et al. Distinct expression patterns of genes associated with muscle growth and adipose deposition in Tibetan pigs: a possible adaptive mechanism for high altitude conditions. High Alt Med Biol 2009;10(1):45–55.
- Ullrich SE, Schmitt DA. The role of cytokines in UVinduced systemic immune suppression. J Dermatol Sci 2000;23(Suppl 1):S10–2.
- Baadsgaard O, Fox DA, Cooper KD. Human epidermal cells from ultraviolet light-exposed skin preferentially activate autoreactive CD4+2H4+ suppressor-inducer lymphocytes and CD8+ suppressor/cytotoxic lymphocytes. J Immunol 1988;140(6):1738–44.
- Dudley AC, Thomas D, Best J, et al. The STATs in cell stresstype responses. Cell Commun Signal 2004;2(1):8.
- 62. Shuai K, Liu B. Regulation of JAK-STAT signalling in the immune system. Nat Rev Immunol 2003;**3**(11):900–11.
- Stempien-Otero A, Karsan A, Cornejo CJ, et al. Mechanisms of hypoxia-induced endothelial cell death. Role of p53 in apoptosis. J Biol Chem 1999;274(12):8039–45.
- 64. Yang W, Qi Y, Lu B, et al. Gene expression variations in highaltitude adaptation: a case study of the Asiatic toad (Bufo gargarizans). BMC Genet 2017;18(1):62.
- Storz JF, Cheviron ZA. Functional genomic insights into regulatory mechanisms of high-altitude adaptation. Adv Exp Med Biol 2016;903:113–28.
- Cheviron ZA, Brumfield RT. Genomic insights into adaptation to high-altitude environments. Heredity (Edinb) 2012;108(4):354–61.
- Bollmer JL, Vargas FH, Parker PG. Low MHC variation in the endangered Galapagos penguin (Spheniscus mendiculus). Immunogenetics 2007;59(7):593–602.
- Wan QH, Zhu L, Wu H, et al. Major histocompatibility complex class II variation in the giant panda (Ailuropoda melanoleuc a). Mol Ecol 2006;15(9):2441–50.
- Zeng QQ, Zhong GH, He K, et al. Molecular characterization of classical and nonclassical MHC class I genes from the golden pheasant (*Chrysolophus pictus*). Int J Immunogenet 2016;43(1):8–17.
- Kan XZ, Yang JK, Li XF, et al. Phylogeny of major lineages of galliform birds (Aves: Galliformes) based on complete mitochondrial genomes. Genet Mol Res 2010;9(3):1625–33.
- Nishibori M, Shimogiri T, Hayashi T, et al. Molecular evidence for hybridization of species in the genus Gallus except for Gallus varius. Anim Genet 2005;36(5):367–75.
- Liu T-K, Chen Y-G, Chen W-S, et al. Rates of cooling and denudation of the Early Penglai Orogeny, Taiwan, as assessed by fission-track constraints. Tectonophysics 2000;**320**(1):69– 82.
- Osozawa S, Shinjo R, Armid A, et al. Palaeogeographic reconstruction of the 1.55 Ma synchronous isolation of the Ryukyu Islands, Japan, and Taiwan and inflow of the Kuroshio warm current. International Geology Review 2012;54(12):1369–88.
- Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 2014;30(15):2114–20.
- Leggett RM, Clavijo BJ, Clissold L, et al. NextClip: an analysis and read preparation tool for Nextera long mate pair libraries. Bioinformatics 2014;30(4):566–8.
- Boetzer M, Henkel CV, Jansen HJ, et al. Scaffolding pre-assembled contigs using SSPACE. Bioinformatics 2011;27(4):578–9.

- Gurevich A, Saveliev V, Vyahhi N, et al. QUAST: quality assessment tool for genome assemblies. Bioinformatics 2013;29(8):1072–5.
- Wood DE, Salzberg SL. Kraken: ultrafast metagenomic sequence classification using exact alignments. Genome Biology 2014;15(3):R46.
- 79. Langmead B, Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nat Methods 2012;9(4):357–9.
- Dobin A, Davis CA, Schlesinger F, et al. STAR: ultrafast universal RNA-seq aligner. Bioinformatics 2013;29(1):15–21.
- Quinlan AR, Hall IM. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 2010;26(6):841–2.
- Krzywinski M, Schein J, Birol I, et al. Circos: an information aesthetic for comparative genomics. Genome Res 2009;19(9):1639–45.
- Slater GS, Birney E. Automated generation of heuristics for biological sequence comparison. BMC Bioinformatics 2005;6:31.
- Kim D, Pertea G, Trapnell C, et al. TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions. Genome Biology 2013;14(4):R36.
- Grabherr MG, Haas BJ, Yassour M, et al. Full-length transcriptome assembly from RNA-Seq data without a reference genome. Nature Biotechnology 2011;29(7):644–52.
- Haas BJ, Delcher AL, Mount SM, et al. Improving the Arabidopsis genome annotation using maximal transcript alignment assemblies. Nucleic Acids Res 2003;31(19):5654– 66.
- Eddy SR. Profile hidden Markov models. Bioinformatics 1998;14(9):755–63.
- Holt C, Yandell M. MAKER2: an annotation pipeline and genome-database management tool for second-generation genome projects. BMC Bioinformatics 2011;12:491.
- Edgar RC. MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res 2004;32(5):1792–7.
- Castresana J. Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. Mol Biol Evol 2000;17(4):540–52.
- Stamatakis A. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 2014;30(9):1312–3.

- Posada, D. (2003) Using MODELTEST and PAUP* to select a model of nucleotide substitution. Curr Protoc Bioinformatics Chapter 6, Unit 6.5 10.1002/0471250953.bi0605s00, 18428705
- Hedges SB, Dudley J, Kumar S. TimeTree: a public knowledge-base of divergence times among organisms. Bioinformatics 2006;22(23):2971–2.
- Bell MA, Lloyd GT. Strap: an R package for plotting phylogenies against stratigraphy and assessing their stratigraphic congruence. Palaeontology 2015;58(2):379–89.
- Han MV, Thomas GW, Lugo-Martinez J, et al. Estimating gene gain and loss rates in the presence of error in genome assembly and annotation using CAFE 3. Mol Biol Evol 2013;30(8):1987–97.
- 96. Yang Z. PAML 4: phylogenetic analysis by maximum likelihood. Mol Biol Evol 2007;**24**(8):1586–91.
- Bleazard T, Lamb JA, Griffiths-Jones S. Bias in microRNA functional enrichment analysis. Bioinformatics 2015;31(10):1592–8.
- Lu TP, Lee CY, Tsai MH, et al. miRSystem: an integrated system for characterizing enriched functions and pathways of microRNA targets. PLoS One 2012;7(8):e42390.
- Bao J, Reecy JM. CateGOrizer: a web-based program to batch analyze gene ontology classification categories. Online Journal of Bioinformatics 2008;9(2):108–12.
- 100. Bindea G, Mlecnik B, Hackl H, et al. ClueGO: a Cytoscape plug-in to decipher functionally grouped gene ontology and pathway annotation networks. Bioinformatics 2009;25(8):1091–3.
- 101. Kanehisa M, Goto S. KEGG: Kyoto encyclopedia of genes and genomes. Nucleic Acids Res 2000;**28**(1):27–30.
- 102. Kearse M, Moir R, Wilson A, et al. Geneious Basic: an integrated and extendable desktop software platform for the organization and analysis of sequence data. Bioinformatics 2012;**28**(12):1647–9.
- 103. Bernt M, Donath A, Juhling F, et al. MITOS: improved de novo metazoan mitochondrial genome annotation. Mol Phylogenet Evol 2013;69(2):313–9.
- 104. Lee CY, Hsieh PH, Chiang LM, et al. Supporting data for "Whole-Genome De Novo Sequencing Reveals Unique Genes that Contributed to the Adaptive Evolution of the Mikado Pheasant." GigaScience Database 2018. http://dx.d oi.org/10.5524/100431.