

Edinburgh Research Explorer

A high-quality genome and comparison of short- versus longread transcriptome of the palaearctic duck Aythya fuligula (tufted duck)

Citation for published version:

Mueller, RC, Ellström, P, Howe, K, Uliano-Silva, M, Kuo, RI, Miedzinska, K, Warr, A, Fedrigo, O, Haase, B, Mountcastle, J, Chow, W, Torrance, J, Wood, JMD, Järhult, JD, Naguib, MM, Olsen, B, Jarvis, ED, Smith, J, Eöry, L & Kraus, RHS 2021, 'A high-quality genome and comparison of short- versus long-read transcriptome of the palaearctic duck Aythya fuligula (tufted duck)', *GigaScience*, vol. 10, no. 12. https://doi.org/10.1093/gigascience/giab081

Digital Object Identifier (DOI):

10.1093/gigascience/giab081

Link:

Link to publication record in Edinburgh Research Explorer

Document Version:

Publisher's PDF, also known as Version of record

Published In:

GigaScience

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.





https://doi.org/10.1093/gigascience/giab081 DATANOTE

DATANOTE

A high-quality genome and comparison of shortversus long-read transcriptome of the palaearctic duck Aythya fuliqula (tufted duck)

Ralf C. Mueller 1,2, Patrik Ellström 3,*, Kerstin Howe 4, Marcela Uliano-Silva 4, Richard I. Kuo 5, Katarzyna Miedzinska 5, Amanda Warr 5, Olivier Fedrigo 6, Bettina Haase 6, Jacquelyn Mountcastle 6, William Chow 4, James Torrance 4, Jonathan M.D. Wood 4, Josef D. Järhult 3, Mahmoud M. Naguib 7, Björn Olsen 3, Erich D. Jarvis 8, Jacqueline Smith 5, Lél Eöry 5 and Robert H.S. Kraus 1,2

¹Department of Migration, Max Planck Institute of Animal Behavior, Radolfzell, 78315, Germany; ²Department of Biology, University of Konstanz, Konstanz, 78457, Germany; ³Department of Medical Sciences, Zoonosis Science Center, Uppsala University, Uppsala, SE-75185, Sweden; ⁴Tree of Life, Wellcome Sanger Institute, Cambridge CB10 1SA, UK; ⁵The Roslin Institute and Royal (Dick) School of Veterinary Studies, University of Edinburgh, Easter Bush, Midlothian EH25 9RG, UK; ⁶Vertebrate Genome Laboratory, The Rockefeller University, New York, 10065, NY; ⁷Department of Medical Biochemistry and Microbiology, Zoonosis Science Center, Uppsala University, Uppsala, 75237, Sweden and ⁸Vertebrate Genome Laboratory and HHMI, The Rockefeller University, New York, 10065, NY

*Correspondence address. Ralf C. Mueller, Am Obstberg 1, 78315, Germany. E-mail: rmueller@ab.mpg.de thttp://orcid.org/0000-0001-8920-6061

Abstract

Background: The tufted duck is a non-model organism that experiences high mortality in highly pathogenic avian influenza outbreaks. It belongs to the same bird family (Anatidae) as the mallard, one of the best-studied natural hosts of low-pathogenic avian influenza viruses. Studies in non-model bird species are crucial to disentangle the role of the host response in avian influenza virus infection in the natural reservoir. Such endeavour requires a high-quality genome assembly and transcriptome. Findings: This study presents the first high-quality, chromosome-level reference genome assembly of the tufted duck using the Vertebrate Genomes Project pipeline. We sequenced RNA (complementary DNA) from brain, ileum, lung, ovary, spleen, and testis using Illumina short-read and Pacific Biosciences long-read sequencing platforms, which were used for annotation. We found 34 autosomes plus Z and W sex chromosomes in the curated genome assembly, with 99.6% of the sequence assigned to chromosomes. Functional annotation revealed 14,099 protein-coding

genes that generate 111,934 transcripts, which implies a mean of 7.9 isoforms per gene. We also identified 246 small RNA families. Conclusions: This annotated genome contributes to continuing research into the host response in avian influenza virus infections in a natural reservoir. Our findings from a comparison between short-read and long-read reference transcriptomics contribute to a deeper understanding of these competing options. In this study, both technologies complemented each other. We expect this annotation to be a foundation for further comparative and evolutionary genomic studies, including many waterfowl relatives with differing susceptibilities to avian influenza viruses.

Keywords: tufted duck; Aythya fuliqula; genome annotation; transcriptome sequencing; Vertebrate Genomes Project; Iso-Seq; Pacific Biosciences; RNA sequencing; small RNA

Background

The tufted duck (Aythya fuliqula, NCBI:txid219594) is a non-model organism that has received attention because of its role in the zoonotic ecology of avian influenza A viruses (AIVs). As a member of the Anatidae family of ducks, it is closely related to the mallard (Anas platyrhynchos), the primary natural host of AIV [1-5]. The Aythya and Anas genera shared a most recent common ancestor ~5 million years ago [6]. However, in contrast to mallards, which carry AIV largely asymptomatically, tufted ducks are less commonly infected with low pathogenic AIV (LPAIV) (see [7] for an updated review on LPAIV infections in tufted ducks) but experience high mortality in outbreaks of highly pathogenic AIV (HPAIV) [8-11]. The tufted duck is a diving duck with a breeding range throughout northern Eurasia, where it is largely a seasonal short-distance migrant. Although it generally feeds in deeper waters than mallards and other dabbling ducks, it generally shares its habitat and roosting areas with these and many other waterfowl species, even leading to a high rate of interspecific hybridization [12, 13]. Hence, given a frequent exposure to AIVs circulating in such habitats, the differences in susceptibility to—and outcome of—AIV infection between these species are likely related to genetic differences affecting, e.g., receptor expression or host response. For example, studies of virus attachment patterns to tissue samples have shown that many AIV subtypes bind less abundantly to intestinal epithelial cells of tufted ducks compared to mallards [7, 14]. In addition, the resistance of mallards against severe HPAIV infection has been partially attributed to the presence of the RIG-Igene and its strong interferon response, in contrast to chickens, which lack this gene and develop severe disease upon HPAIV infection [15]. Future studies in non-model bird species such as the tufted duck are important to disentangle the role of the host response and other genetic factors in AIV infection and aid in our understanding of the zoonotic ecology of AIV in the natural reservoir. A prerequisite for such studies is a well-assembled and annotated genome and transcriptome [16].

Developments in omic sequencing technologies over the past 2 decades have revolutionized biology. Instead of studying single genes and their products, whole genomes and transcriptomes can now be readily sequenced and assembled at a lower cost than before. Massive parallelization and high throughput in next-generation sequencing (NGS) have decreased sequencing costs and ultimately increased sequencing depth [17]. This allows for whole-genome sequencing of any species and opens up new possibilities for in-depth studies related to infection biology and host response to external stressors beyond model species in which a rich genetic toolbox can be deployed [18, 19]. Nonmodel species are frequently understudied, yet they are exposed to environmental stressors such as infectious diseases, which they can transmit to livestock and humans [20]. Approximately 70% of human infectious diseases are zoonoses [21, 22]. An indepth understanding of a pathogen's zoonotic ecology in the animal reservoir is important to prevent human infections. This, in turn, requires studies of host-pathogen interactions at the genetic level. NGS helps bridge the gap between model and nonmodel species [23], and with third-generation long-read-based sequencing, high-quality reference genomes are now also available for non-model organisms, as in the Vertebrate Genomes Project (VGP) [24]. This is supplemented by affordable long-read RNA sequencing, making de novo assembled transcriptomes unnecessary [25].

Transcriptome annotation of the genome used to be constrained to either low-throughput and costly complementary DNA (cDNA) cloning or Illumina's high-throughput short-read RNA sequencing (RNA-Seq). High-quality short-read RNA sequencing combined with a reference genome allows for a reasonable transcriptome reconstruction. However, there are some caveats: Owing to alternative splicing, a single gene can have multiple alternative variants (isoforms) and as a consequence can be translated into proteins with different functions [26, 27]. Illumina short reads need to be assembled into transcripts, which can lead to incompleteness and errors in transcript model reconstruction. This biases the correct inference of isoforms and thus misses the transcriptome's complexity [26, 28]. Furthermore, because short-read sequencing is also limited in GC-rich regions and regions of low sequence complexity [29], not all transcripts are recovered. In contrast, in full-length transcript isoform sequencing (e.g., Pacific Biosciences [PacBio] Iso-Seq) of messenger RNA, the single direct reads from 5' to 3' usually do not need to be assembled and thus prevent assembly ambiguities, which facilitates the detection of novel isoforms [30, 31] and accurate reconstruction of transcript structure [29]. This thirdgeneration transcript sequencing technology also allows for a much more detailed functional annotation. It is crucial to recover as many isoforms as possible for functional studies [32-34]. For example, the immunome comprises a set of genes associated with immune processes and is a heavily complex portion of the genome [25, 35]. Thus, the benefits of third-generation sequencing must be exploited to their fullest potential to facilitate studies in bird immunogenomics [36] and improve databases such as the Avian Immunome DB for comparative analyses in non-model bird species [37].

Data Description

This study presents the first high-quality, chromosome-level reference genome assembly of the tufted duck using the VGP pipeline [24] followed by manual curation [38], which we annotated with short- and long-read transcripts from 6 different tissues, and short reads from small RNAs from the same tissues. We further contrasted different sequencing technologies and bioinformatics pipelines, and tissue-specific comparisons of expressed genes. This annotation serves as a foundation for

Table 1: Assembly statistics of the tufted duck genome

Statistic	Value
Genome coverage (x)	64.03
Total sequence length (bp)	1,127,004,725
Ungapped sequence length (bp)	1,117,587,328
No. of scaffolds	105
Scaffolds assigned to	36 + 1 mitochondrion
chromosomes	
Unplaced scaffolds	68
Contig NG50 (bp)	17,816,505
Scaffold NG50 (bp)	85,905,639
Base-call error in 10 kb	<1 nucleotide

further comparative and evolutionary genomics, and gene expression experiments in the tufted duck and its many waterfowl relatives with different susceptibilities to AIV [36, 39].

Analyses

Reference genome assembly

The reference genome was constructed according to VGP's 6.7.P2.Q40.C99 metric standards [24]. This includes contig NG50 $> 10^6$ bp, scaffold NG50 $> 10^7$ bp, most of the genome separated into haplotypes (P2), Phred-scaled base accuracy > Q40, and 99.5% of the assembly assigned to chromosomes. The contig NG50 of the tufted duck genome assembly was 17.8 Mb. The curation (as in Howe et al. [38]) resulted in 34 removals of misjoins, 34 joins previously missed, and 4 removals of false duplications. This reduced the primary assembly length by 0.8% and increased the scaffold NG50 by 18.7% to 85.9 Mb whilst decreasing the scaffold number from 123 to 105. The total sequence length of the assembly after curation was 1.127 Gb, which is close to the expected size of duck genomes (1.25-1.34 Gb) [40]. During curation, telomeres were not detected, and a sweep for centromeres with sequences from [41] revealed no results. Chromosomalscale units were identified and named by size. Of the expected 39 chromosome pairs according to the karyotype [42], 34 autosomes plus Z and W could be identified, with 99.6% of the sequence assigned to them (Table 1, Fig. 1). The pseudoautosomal region in W/Z (\sim 2 Mb) collapsed into a single copy on the Z chromosome (Supplementary Fig. S1). Detailed assembly statistics by chromosome can be found Supplementary Table S1.

The raw data have been deposited in the GenomeArk repository [43]. The curated primary assembly has been deposited in NCBI under accession No. GCF_009819795.1 [44] and can be browsed in the Genome Data Viewer [45]. A comparison of assembly metrics before and after curation can be found in Supplementary Table S1.

Analysis of repetitive sequences

Repetitive element content and composition in the tufted duck genome assembly was identified and compared with that of the mallard genome (ZJU1.0). Overall, 13% and 15% of the tufted duck and mallard genomes, respectively, are made up by repeats. The most abundant repeat classes are long interspersed nuclear elements followed by long terminal repeats, short interspersed nuclear elements, and DNA repeats. The genomic repeat composition, in general, is similar between tufted duck and mallard, although the mallard genome has slightly higher repeat content. The only major difference we observed was in repeats that were not classified by RepeatModeler into any known repeat categories. These are termed "Unknown," and the mallard genome seems to contain 4.4 times as many Unknown repeats as the tufted duck genome (Supplementary Fig. S2). Nevertheless, when compared at chromosome level (Supplementary Fig. S3), repeat content is similar between the orthologous pairs, including autosomes and sex chromosomes, and the major difference is only observed in unplaced scaffolds. Forty percent of the mallard unplaced scaffolds are made up by Unknown repeats, while it is only 6% in the tufted duck.

Thirteen telomeric and 3 potential centromeric repeat regions were identified in the tufted duck genome (Supplementary Fig. S4).

Gene/transcript model reconstruction with Illumina RNA-Seq and PacBio Iso-Seq reads

With the Illumina RNA-Seq, a mean (SD) of 97.65% (0.51%) of the reads were retained after adapter and quality trimming. For the PacBio Iso-Seg, a mean (SD) of 80.57% (3.84%) full-length nonchimeric (FLNC) eads were retained after error correction (Table 2).

HISAT2 could map a mean (SD) of 93.21% (0.64%) Illumina RNA-Seg short reads and Minimap2 could map 97.39% (1.34%) PacBio Iso-Seq long reads to the reference genome. The mean (SD) read length of the long-read data was 1,214.0 (262.0) nt, an order of magnitude longer than that of the short reads (131.5 [0.7] nt). However, the mean number of reads was almost 600fold higher with short-read data (129,411,992 [11,989,532]) compared to long-read data (217,293 [143,219]), implying a 60 times higher sequencing depth with the short reads. Not surprisingly, StringTie2 (in the short-read pipeline) assembled more transcript models and inferred more genes and exons than the longread pipeline. This was true for all tissues except in the lung; more transcript models were inferred in the long-read pipeline. Lung RNA (cDNA) was sequenced on 2 Zero-mode waveguides (ZMW) and produced the highest numbers of subreads and FLNC after processing. The highest number of genes in each pipeline was predicted for ovary (Illumina) and brain (PacBio). The highest number of transcript models was predicted for ovary (Illumina) and lung (PacBio). The same pattern applies to predicted exons (Table 3).

In the Illumina short-read RNA-Seq transcript model reconstruction, more exons per gene were recovered on average than in the long-read pipeline (Table 3). The distribution of exons per gene in both pipelines generally followed the same pattern, with a decreasing number of genes as the exons per gene increased. However, there is a remarkable difference in the short-read data, which recovered more 2-exon genes than single-exon genes for all tissues (Fig. 2). This pattern is even more pronounced in the exons per transcript analysis (Supplementary Fig. S5).

In the PacBio long-read Iso-Seq transcript model reconstruction, more transcripts per gene were recovered on average than in the short-read pipeline (Table 3). The distribution of transcripts per gene followed the same pattern, with a decreasing number of genes as the transcripts per gene increased. However, these numbers had a slower decay in the long-read data, indicating a better recovery of multi-transcript genes (Fig. 3).

Functional annotation of merged transcripts

After merging all 12 transcript model reconstructions (6 tissues with short reads and long reads), 345,870 transcripts with 2,381,662 exons were predicted in 49,746 genes (Table 4). Of these, 178,198 transcripts (or 16,758 genes) were predicted by

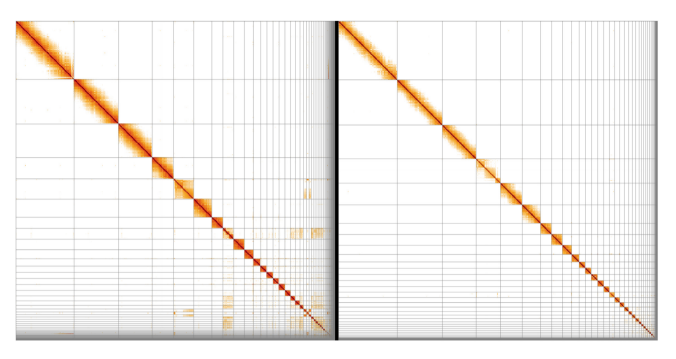


Figure 1: HiGlass Hi-C 2D maps of the tufted duck genome assembly before (left) and after (right) manual curation. Off-diagonal hits indicate missing joins, which have been corrected during curation. Broken patterns within scaffolds (e.g., at the end of the first scaffold before curation) can indicate intra-scaffold misassemblies, which were also addressed during curation. They can, however, also be features of the respective chromosome, as in the fourth post-curation scaffold, the structure of which was corrected and asserted during curation.

Table 2: Illumina RNA-Seq reads before and after trimming, PacBio Iso-Seq reads before and after error correction, ZMW yield, and the number of FLNC reads

Platform	Parameter	Brain	Ileum	Lung	Ovary	Spleen	Testis
Illumina	No. raw reads (PE)	71,648,303	72,410,462	63,191,981	72,442,392	68,315,029	53,747,262
	No. trimmed (paired)	70,025,835	70,902,250	61,059,701	70,892,177	66,813,691	52,652,923
PacBio	No. subreads	9,249,099	12,363,369	19,206,097	1,279,561	8,401,115	12,616,953
	No. CCS	158,698	415,314	529,108	68,112	167,077	288,984
	ZMW yield (%)	15.87	41.53	26.46	3.41	16.71	28.90
	Mean No. of passes	58.3	29.8	36.3	18.8	50.3	43.7
	No. FLNC	133,684	343,634	432,817	49,887	134,124	234,423

Two ZMW were used for lung and ovary. CCS: circular consensus sequencing; FLNC: full-length, non-chimeric; PE: paired end; ZMW: zero-mode waveguide.

Table 3: Transcript model reconstruction per tissue and pipeline

Parameter	Platform	Brain	Ileum	Lung	Ovary	Spleen	Testis
Mapped (%)	Illumina	92.92	92.98	93.42	94.33	92.44	93.19
	PacBio	96.66	98.13	97.92	99.40	95.67	96.57
No. genes	Illumina	22,348	20,838	20,692	32,046	21,608	29,225
	PacBio	15,776	10,813	12,912	8,862	6,773	11,746
No. transcripts	Illumina	44,808	40,968	43,877	77,997	42,000	57,758
	PacBio	37,601	35,284	46,587	19,513	14,030	28,852
No. exons	Illumina	422,741	395,719	412,225	569,566	383,679	483,444
	PacBio	138,038	217,391	243,566	119,080	71,211	160,380
Exons per gene (mean)	Illumina	18.9	19.0	19.9	17.8	17.8	16.5
	PacBio	8.7	20.1	18.9	13.4	10.5	13.7
Transcripts per gene (mean)	Illumina	2.0	2.0	2.1	2.4	1.9	2.0
	PacBio	2.4	3.3	3.6	2.2	2.1	2.5

CPC2 to be protein-coding. UniProt hits were found for 208,274 transcripts (or 17,911 protein-coding genes). Of these, 4,766 genes had no long-read support and 432 genes no short-read support in the data. The number of protein-coding genes pre-

dicted by CPC2 and number of hits in UniProt overlapped for 15,103 genes (Supplementary Fig. S6). Conservative filtering of the annotation (ignoring features flagged as nonsensemediated decay and only keeping full-length hits in UniRef50

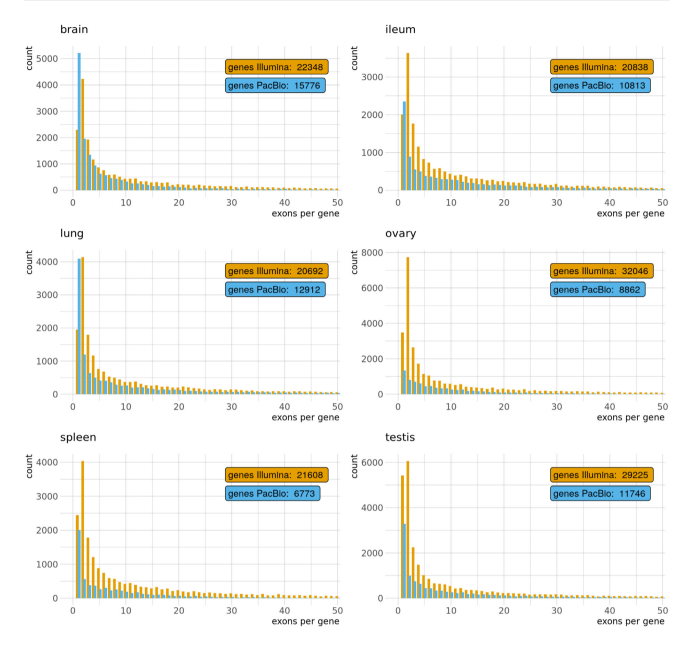


Figure 2: Distribution of single- and multi-exon genes per tissue and pipeline. Only the first 50 groups are shown.

Table 4: Functional annotation categorized by different matches

	_	-	
Parameter	Genes	Transcripts	Isoforms
Total No. of entries	49,746	345,870	7.0
UniRef50 total hits	17,911	208,274	11.6
UniRef50 match			
Full	13,024	99,737	7.7
90%	4,937	12,197	2.5
50%	3,208	6,540	2.0
<50%	2,474	4,447	1.8
≥50%	14,731	118,474	8.0
≥90%	14,099	111,934	7.9
No hit (but full-length)	27,787	78,860	

Note: UniRef50 total hits also includes 5' degraded records, whereas the match classes only include full-length records.

that matched with ≥90%) resulted in 111,934 transcripts and 14,099 protein-coding genes. This result implies a mean of 7.9 isoforms per gene (Table 4). Predicted and annotated genes by chromosome can be found in Supplementary Table S2.

Of 78,860 full-length transcripts with no UniRef50 hit, 62,147 were flagged as potentially protein-coding (and the remainder as nonsense-mediated decay), and 26,489 as single-exonic while the remainder as composed of ≥ 2 exons.

RIG-I/DDX58 is intact and expressed in tufted duck

In the mallard genome [46, 47], RIG-I/DDX58 is annotated on chromosome Z (NC_051804.1), position 69,037,671-69,061,400 (23,730 nt), and consists of 18 exons. Searching the protein sequence in the tufted duck genome assembly produced 1 significant alignment with the predicted gene XM_032205362.1, also on chromosome Z (max score: 1,882, total score: 1,882, query

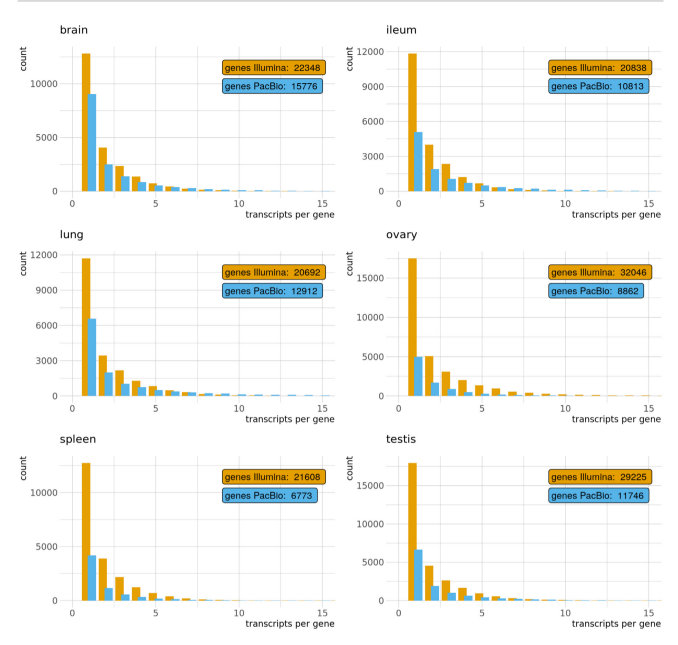


Figure 3: Distribution of single- and multi-transcript genes per tissue and pipeline. Only the first 15 groups are shown.

cover: 100%, E-value: 0.0, percent identity: 97.00%; see alignment in Supplementary Listing 1).

Searching the predicted open reading frames (ORFs) from the functional annotation in the 2 mallard RIG-I/DDX58 isoforms revealed 9 matches after conservative filtering (≥90% match; Table 5). Gene G24916 on chromosome 6 (NC.045564.1) matched with ORFs of 6 transcripts while G24916.1 contained the same translated amino acid sequence as G24916.2 and G24916.4. Gene G46857 on chromosome Z (NC.045593.1) matched with ORFs of 3 transcripts while G46857.2 contained the same translated amino acid sequence as G46857.3. ORFs G24916.7/8 and G46857.2/3/4 were flagged with "5prime_degrade," which means that the transcript might be incomplete on the 5′ end. Both genes were predicted in the short-read and long-read pipeline. A detailed list of reconstructed transcript models by pipeline and tissue for these 2 genes can be found in Supplementary Table S3.

All G24916 sequences relate to the IFIH1 gene (encodes MDA5, a RIG-I-like receptor) while G46857 sequences relate to the RIG-I/DDX58 gene. An alignment of the mallard and tufted duck RIG-I/DDX58 amino acid sequences revealed 15 variants (14 substitutions, 1 insertion), which were predicted to have no effect on the biological function of the protein (Supplementary Tables S4 and S5). The nucleotide sequences 1 kb upstream of the RIG-I/DDX58 gene contained 4 identical transcription factor binding sites (Nkx2-5, NF-kappB p65, c-Rel, NF-kappaB) in each species (Supplementary Table S6).

Tissue-specific expression and intersections

Of the 17,911 genes in UniRef50 (total hits), 4,766 were exclusively supported by short reads and 432 by long reads. In the short-read pipeline, 11,165 genes intersected across all tissues. The highest number of exclusively expressed genes was in testis

Table 5: Blastp results (≥90% match) of predicted ORFs from the functional annotation searched in 2 mallard RIG-I/DDX58 isoforms NP_001297309.1 (933 aa) and XP_038025643.1 (988 aa)

Mallard	Tufted duck								
Isoform	Chromosome	ORF	Start/End	nt	Frame	Strand	Exons	aa	
NP_001297309.1	NC_045564.1 (6)	G24916.1	21,885,153/21,914,958	29,805	F2	+	17	1,003	
NP_001297309.1	NC_045564.1 (6)	G24916.2	21,885,153/21,914,958	29,805	F1	+	16	1,003	
NP_001297309.1	NC_045564.1 (6)	G24916.3	21,887,313/21,915,355	28,042	F1	+	17	994	
NP_001297309.1	NC_045564.1 (6)	G24916.4	21,887,313/21,914,044	26,731	F1	+	16	1,003	
XP_038025643.1	NC_045564.1 (6)	G24916.7*	21,887,508/21,915,353	27,845	F1	+	16	1,044	
XP_038025643.1	NC_045564.1 (6)	G24916.8*	21,887,508/21,915,503	27,995	F1	+	16	1,040	
NP_001297309.1	NC ₋ 045593.1 (Z)	G46857.2*	69,123,499/69,145,704	22,205	F3	+	18	948	
NP_001297309.1	NC_045593.1 (Z)	G46857.3*	69,123,499/69,147,273	23,774	F3	+	18	948	
NP_001297309.1	NC ₋ 045593.1 (Z)	G46857.4*	69,123,529/69,145,281	21,752	F3	+	18	938	

ORFs marked with an asterisk were flagged with "5prime_degrade," which means that the start codon was not found in the TAMA ORF/NMD prediction pipeline. aa: amino acid.

(988), followed by ovary, brain, spleen, ileum, and lung (Fig. 4). In the long-read pipeline, 2,475 genes intersected across all tissues. The highest number of exclusively expressed genes was in brain (779), followed by testis, ileum, lung, ovary, and spleen (Fig. 5). Overlap of expressed genes from each pipeline can be found in Supplementary Fig. S7.

Small RNA analyses

For the TruSeq small RNA sequencing data, a mean (SD) of 95.81% (3.51%) of the reads were retained after adapter and quality trimming (Table 6). Overall, STAR could map a mean (SD) of 93.91% (5.28%) of these reads to the reference genome, which divides into 67.99% (13.75%) uniquely mapped reads and 25.92% (12.08%) multi-mapped reads (\leq 10 loci). Cufflinks predicted the highest number of genes in the spleen (33,133) followed by testis (31,205). The remaining tissues had much lower numbers of genes, ranging from 8,441 (ileum) to 13,606 (brain). The same pattern applies to the number of predicted transcripts and exons (Table 6).

Each transcript was composed of a mean (SD) of 1.3 (0.2) exons. The distribution of single-exon and multi-exon transcripts shows a clear trend towards single-exon transcripts and a quickly diminishing number of multi-exon transcripts. However, this was less pronounced in spleen but even more so in testis (Fig. 6). The generally decaying length distribution of transcripts shows 2 clear peaks at 20 and 50 bp except for testis with the first peak at 30 bp. Except for spleen and testis data, there are very few transcripts >200 bp (Supplementary Fig. S8).

Scanning the genome (in silico) for Rfam's covariance models of RNAs resulted in 1,234 hits. The same scan on the assembled small RNA transcripts (in vitro) revealed a mean (SD) of 346.5 (26.9) hits across all tissues. After removing lower-scoring overlaps and hits with E-value >5.0E-4 from the cmscan result, 1,076 distinct features were predicted in the tufted duck genome. In the tissue's small RNA assemblies, a mean of 327.5 (26.5) features were annotated, with the same filtering (Table 7). A mean (SD) 93.25% (1.09%) of the in vitro annotated features were predicted by the in silico scan. Furthermore, a mean (SD) of 22.2 (2.3) features were annotated that were not predicted in silico (Table 7).

After further filtering for unique RNA families (Rfam accession numbers/covariance models), 306 distinct RNA families were predicted in the genome, with 246 annotated in all tissues (pooled). The number of predicted and annotated covariance models overlapped for 237 RNA families, while 69 were only

identified in the genome scan and 9 were only identified in the pooled tissue annotations.

Discussion

In this study, we present the first chromosome-level reference genome assembly of the tufted duck. The genome contiguity is on par with other reference bird genome assemblies that used long reads such as mallard [46,47], chicken [48], and recent VGP-pipeline-generated zebra finch [49]. The assembly's contig NG50 of 17.8 Mb is comparable to chicken (17.7 Mb) but considerably higher than in mallard (5.7 Mb) and zebra finch (4.4 Mb). The assembly's scaffold NG50 of 85.9 Mb is higher than in mallard (76.3 Mb) and zebra finch (70.9 Mb) and considerably higher than in chicken (20.8 Mb). All our mapping results from the Illumina short-read and PacBio long-read RNA pipelines confirm full adherence to the VGP 6.7.P2.Q40.C99 standard, which also implies transcript mappability >80% [24].

The higher numbers of recovered genes, transcripts, and exons in the short-read transcript model reconstruction can be mainly explained by the higher sequencing depth and further reinforced by the different RNA preparation protocol. With Illumina, virtually all trimmed, paired-end reads were kept for mapping to the genome while with PacBio, only 5' cap-selected and FLNC reads were kept. However, the transcript model reconstruction in the long-read pipeline often almost matched or even exceeded (lung) the mean number of transcripts in the short-read pipeline. Furthermore, the long-read pipeline recovered more transcripts per gene (isoforms) on average. Taken together, this is remarkable considering the 60-fold higher sequencing depth in the short-read pipeline. It also corroborates the strength of PacBio Iso-Seq, which seems to better reflect the complexity of the transcriptome, considering that transcripts did not need to be assembled but were sequenced full-length. However, this result is not reflected in the functional annotation and, together with the 27,787 putatively intact genes without a hit in the UniRef50 database, may indicate potentially undescribed genes.

In the short-read transcript model reconstruction, more 2-exon genes than single-exon genes were predicted for all tissues, and it seems as if some transcripts could not be assembled entirely or StringTie2 tried to "avoid" single-exon genes. Transcript model reconstruction in StringTie2 is based on the concept of extending short reads to create so-called super-reads [50], which seems appropriate for whole-genome assemblies. In

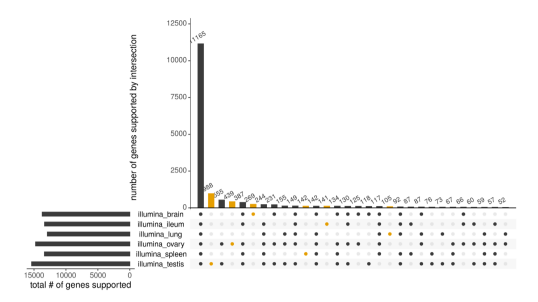


Figure 4: In the short-read data set, the highest total number of supported genes was found in testis (left panel, bottom), followed by ovary, brain, spleen, ileum, and lung. All 6 tissues intersected in 11,165 genes (main panel, left). The highest number of exclusively supported genes was also found in testis (988), and followed the same order as the total number of genes (main panel, yellow).

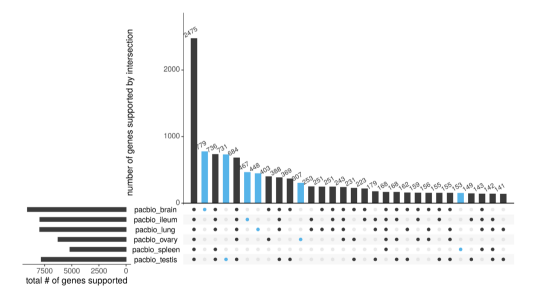


Figure 5: In the long-read data set, the highest total number of supported genes was found in brain (left panel, top), followed by lung, ileum, testis, ovary, and spleen. All 6 tissues intersected in 2,475 genes (main panel, left). The highest number of exclusively supported genes was found in brain (779), followed by testis, ileum, lung, ovary, and spleen (main panel, blue).

Table 6: Small RNA read processing and assembly statistics

Statistic	Brain	Ileum	Lung	Ovary	Spleen	Testis
No. raw reads (PE)	78,078,195	58,021,264	70,381,224	79,425,103	65,767,638	67,436,837
No. trimmed reads (PE)	73,753,404	57,021,189	69,326,112	77,333,115	58,716,681	65,395,835
Mapped uniquely (%)	51.82	88.04	72.04	71.01	72.43	52.59
Mapped multiply (%)	44.90	9.48	24.42	26.59	18.18	31.97
No. genes	13,606	8,441	11,899	9,903	33,133	31,205
No. transcripts	13,685	8,520	11,995	9,954	33,761	31,342
No. exons	17,276	12,650	15,397	11,588	54,504	35,026

PE: paired end.

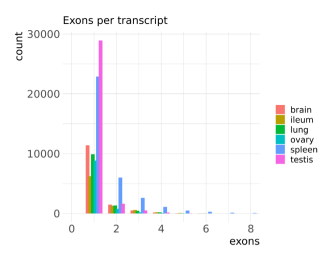


Figure 6: Distribution of single-exon and multi-exon small RNA transcripts for each tissue.

Table 7: Results of cmscan on assembled small RNA transcripts after filtering

Parameter	Brain	Ileum	Lung	Ovary	Spleen	Testis
In vitro	328	294	317	312	345	369
Intersection	310	274	295	293	315	345
Additional	18	20	22	19	30	24

Intersection refers to small RNAs predicted by the *in silico* genome scan. Additional refers to annotated small RNAs that were not detected by cmscan in the reference genome.

transcriptomics, however, multiple splice variants are possible, and with the super-read concept in mind, it may thus be possible that StringTie2 discards a single-exon transcript model in favour of an alternative multi-exon splice variant containing the same exon. This, in turn, might have a substantial impact on the functional annotation based on transcript models solely assembled with short reads. Real single-exon transcripts might be missed, or even worse, false-positive multi-exon transcript models might be reconstructed. We merged transcript models of both pipelines with higher priority on transcript end sites for the long-read-inferred transcript models to mitigate this effect.

While both the number and quality of published vertebrate genome assemblies are increasing, hardly any are complemented by a transcriptome of multiple tissues from the same individual [26, 51]. Automated annotations (e.g., the NCBI Eukaryotic Genome Annotation Pipeline [52]) provide reasonable in silico predictions of coding potential; however, RNA (cDNA) sequencing adds evidence for expressed genes. Based on the inferred transcripts in this study, a total of 14,099 protein-coding genes could be identified in the UniRef50 database after conservative filtering (≥90% match), which is comparable to NCBI's in silico prediction of 15,578 protein-coding genes [53]. The number of identified protein-coding genes in tufted duck is also in line with the prediction in other bird species such as mallard (16,836 [54]), chicken (17,477 [55]), or zebra finch (16,197 [56]). CPC2 predicted 84.2% of the potentially protein-coding genes found in UniRef50, which would serve as a conservative estimate of the proteincoding potential by just looking at the reference genome. However, beyond the 17,911 genes annotated by UniRef50, the annotation contains an additional 27,787 genes with protein-coding potential according to the TAMA ORF/NMD prediction pipeline,

with these being potential candidates for further analyses. Gene and transcript identification in non-model species relies on annotations of preferably closely related model organisms. However, protein-coding genes are mainly described by a single transcript and predominantly built on short-read and comparative data [26].

We could confirm that the gene RIG-I/DDX58 is intact and expressed in the tufted duck, and almost identical and at the same position as in mallard. The differences in amino acid sequence are tolerated, and transcription factor binding sites are identical with those in mallard. Taken together, there are no obvious differences in the RIG-I gene that can account for the difference in susceptibility to influenza seen in each species. This is remarkable considering that tufted ducks are highly susceptible to AIV and indicates that the host response is complex and depends on more than an intact and expressed RIG-I/DDX58 gene [36, 39].

Besides a high-quality transcriptome for the tufted duck, this study also provides a tissue-specific expression atlas. In the short-read pipeline, there is a large decrease in numbers of genes expressed in all tissues to genes exclusively expressed in a single tissue or a few tissues. This distribution is much more balanced in the long-read pipeline and may indicate that the coverage in PacBio was too low to fully recover all genes in all tissues.

The number of 306 Rfam small RNA families predicted (with 246 annotated) for the tufted duck in this study is comparable to 352 families predicted in chicken (Gallus gallus, version 5) in Rfam 14.4 [57]. The peaks of transcript length at 20 and 50 bp are in the area of microRNA (miRNA), which are usually 18-23 nt [58-60], and pre-miRNA, which are in the range of 55-70 nt. The substantial variance across tissues in our data (many more genes, transcripts, and exons in spleen and testis than in the other tissues) might be explained by Illumina sequencing bias introduced at the adapter-ligation step of cDNA library constructions [61, 62]. Furthermore, according to Illumina's TruSeq Small RNA library preparation protocol, small RNA populations can vary significantly between different tissue types and species, and types and coverage vary depending on which bands are selected during gel excision. This is additional support of a strategy to sequence multiple tissues to obtain a fuller picture of small RNA expression in an organism. Gene duplications might explain the significant difference between unique mappings and multiple mappings of small RNA to the genome across tissues. The in silico scan of small RNA in the genome could predict almost all small RNAs in the assembled transcripts. More importantly, however, 22.2 additional small RNAs were discovered on average in the in vitro scan that would otherwise have been unnoticed. Small (non-coding) RNAs play an essential role in gene regulation, translation, and chromosome structure [63, 64] and are often associated with diseases [65, 66]. This vital fraction of the genome is rarely validated in vitro in the genome and transcriptome annotation literature. However, small RNA studies have been continuously increasing over the past 20 years from 1,966 publications in 1999 to 8,034 in 2019 (searching for "small RNA" on Web of Science [67]). Relying on in silico prediction of small RNA alone can lead to misinterpretation of pathways and gene regulation, and sequencing small RNA of nonmodel organisms is, therefore, an advancement for genome annotation [68]. We strongly encourage in vitro sequencing of small RNA in de novo genome and transcriptome studies, or otherwise, the scientific community will miss much detail that will be important to decipher relevant differences in the biology between species.

Potential Implications

This study presents the first high-quality reference genome assembly of the non-model tufted duck species. It is complemented by coding and small non-coding RNA transcriptome annotation from 6 different tissues. The genome assembly contributes to the VGP's ongoing mission to generate near error-free and complete genome assemblies of all extant vertebrate species. By utilizing, comparing, and combining the strengths of low error rates and high sequencing depth in Illumina RNA sequencing, and the full-length transcript sequencing in PacBio's Iso-Seq, this annotation culminates in a merged transcriptome with functional annotation and an expression atlas. Evidence from small RNA of the same tissues sequenced using the Illumina platform revealed small RNAs that would have otherwise remained undetected. Our findings from a comparison between short-read and long-read reference transcriptomics contribute to a deeper understanding of these competing options. In this study, both technologies complemented each other. While short-read data were sufficient to annotate proteincoding genes, long-read data recovered more transcripts per gene and potentially further protein-coding genes that could not be annotated. With the ongoing improvement of base call quality in long-read sequencing, short-read transcriptome sequencing might become expendable, and we recommend reconstructing transcriptomes using long-read sequencing with high coverage. Together, the genome and transcriptome annotation of the tufted duck is an excellent resource for public omics databases and a foundation for downstream studies, e.g., regarding disease response. The data set's high quality for a non-model species allows for a much finer resolution of genetic differences and commonalities in closely related species, which is crucial while studying the reservoirs of zoonotic pathogens.

Methods

Sampling and dissection of tissues

Captive-bred, wild tufted ducks were kept at the animal breeding facility (Swedish National Veterinary Institute, Uppsala, Sweden). The ducks were obtained from Snavelhof breeding farm, Veeningen, the Netherlands, in May 2017. Tissue samples were obtained from 5 females and 5 males (12 months old) after euthanasia with an injection of 1 mL of pentobarbital (100 mg) in the wing vein. The following tissues were collected from the birds: brain, ileum, spleen, lung, and gonads (ovary or testis). Tissues were immediately snap-frozen in liquid nitrogen and stored at -80°C until shipment on dry ice to the Roslin Institute, Edinburgh, UK. All animal experiments were carried out in strict accordance with a protocol legally approved by the regional board of the Uppsala animal ethics committee, Sweden (permission No. 5.8.18-07998/2017). The animal experiments were conducted in biosafety level 2 animal facilities at the Swedish National Veterinary Institute.

Genomic DNA: library preparation, sequencing, and assembly

To obtain both sex chromosomes, DNA was extracted from lung tissue of a female tufted duck. Library preparation and sequencing was conducted as in [24], using 4 types of sequencing data and the VGP assembly pipeline 1.6 (all details given in Rhie et al. [24] and pipeline available on [69]). The sequence data consisted of PacBio chromium linked reads (CLRs) (64.03× coverage), 10X Genomics CLRs (110.83× coverage), Bionano Genomics optical maps created by direct labelling and staining (DLS; 371.51× coverage), and chromatin conformation capture coupled with high-throughput sequencing (Hi-C; 92.27× coverage) (Arima Genomics, San Diego, CA, USA). In brief, PacBio reads were input to the diploid-aware long-read assembler FALCON and its haplotype-resolving tool FALCON-Unzip [70]. The resulting primary contigs were input to the Purge-Dups pipeline [71] to identify and remove remaining haplotigs in the primary set. In the next step, primary-purged contigs were subject to 2 rounds of scaffolding using the 10X long molecule linked reads. Further, pre-assembled DLS Bionano cmaps were applied for further scaffolding and ordering using the Solve pipeline (Bionano Genomics, San Diego, CA, USA). The resulting scaffolds were then further scaffolded into chromosome-scale scaffolds using Hi-C data and the Salsa2 pipeline [72]. Finally, the primary assembly plus the Falcon-phased haplotigs were concatenated for 3 rounds of base call polishing: first with PacBio reads and Arrow software [73] and subsequently 2 rounds of polishing with 10X linked reads and FreeBayes software (FreeBayes, RRID:SCR_01076 1) [74]. The genome was decontaminated and manually curated as described in Howe et al. [38].

Tissue preparation, DNA and RNA extraction

For disruption and homogenization of tissues, snap-frozen samples were ground to a fine powder under liquid nitrogen using a mortar and pestle. Samples were transferred to 1.5 mL frozen tubes and kept on dry ice until further processing. Total RNA was obtained following a standard TRIzol protocol with DNase treatment and column purification. Small RNA was prepared according to the miRNeasy kit protocol 217004 (Qiagen, Venlo, Netherlands). Integrity and quality of the RNA were confirmed by electrophoresis on an Agilent 2200 Tapestation using appropriate screen tapes. The concentration was determined using the Nanodrop ND-1000 (Thermo Fisher Scientific, Waltham, MA, USA) (Supplementary Tables S7 and S8). For DNA extraction and sequencing, powdered lung tissue was sent to the Vertebrate Genomes Lab (VGL) at Rockefeller University, New York, NY, USA.

Illumina cDNA library preparation and sequencing

RNA was sent to Edinburgh Genomics, Edinburgh, UK, for library preparation and sequencing on an Illumina HiSeq 4000 platform (Illumina HiSeq 3000/HiSeq 4000 System, RRID:SCR_016386) [75] with 2 \times 150 bp paired-end reads using the TruSeq library preparation protocol (stranded). Median (SD) insert size was 137-148 (67–81 bp), yielding \geq 290 M + 290 M reads per sample. Small RNA was also sent to Edinburgh Genomics for TruSeq small RNA library preparation and sequencing using a NovaSeq 6000 platform (Illumina NovaSeq 6000 Sequencing System, RRID:SCR_016 387) [76] with 2×50 bp paired-end reads. Median insert size was 141–144 bp, yielding \geq 225 M + 225 M reads per sample.

Genome analysis and comparison with the mallard genome assembly

Repeat content in the tufted duck genome assembly was defined using RepeatMasker v4.1.0 (RepeatMasker, RRID:SCR_01295 4) [77] with duck-specific repeat sequences from the combined Dfam v3.1 (Dfam, RRID:SCR_021168) [78] and RepBase v20170127 (Repbase, RRID:SCR_021169) [79] repeat libraries. RepeatMasker was run in "sensitive" mode (-s) using "Aythya fuligula" as the query species (-species 'Aythya fuligula'). This was followed by a second round of repeat masking, which was carried out using a

novel repeat sequence library obtained by RepeatModeler v2.0.1 (RepeatModeler, RRID:SCR_015027) [80]. To generate a comparative data set on the mallard genome (ZJU1.0 [46, 47]) we used the same repeat-masking strategy.

Telomeric repeats were identified by searching for known vertebrate-specific repeat hexamers of "TTAGGG" and "CCC-TAA," while known Anseriformes-specific centromeric repeat sequences [81] were mapped with RepeatMasker.

Orthologous chromosome pairs were identified by searching for synteny between the tufted duck and mallard genomes. The 2 genomes were aligned with Minimap2 (Minimap2, RRID: SCR_018550) [82] using options "-secondary=no -asm 10" and the primary alignments were used as a proxy for synteny between the 2 genomes. Primary alignments between regions are shown in a circos plot [83] in Supplementary Fig. S9.

PacBio cDNA library preparation and sequencing

Two micrograms of total RNA from each sample in 4 parallel reactions were converted to cDNA using the Teloprime full-length cDNA amplification kit (013, v1) according to the manufacturer's instructions (Lexogen, Vienna, Austria). After end-point PCRs, all samples were tested for quality and quantity. The product size distribution was visualized using an Agilent 2200 Tapestation using D5000 screen tape. The library concentration was measured on a Qubit 3 (Thermo Fisher Scientific, Waltham, MA, USA) with high-sensitivity DNA reagents (Supplementary Table S9). Technical replicates were pooled and selected for PacBio Iso-Seq assays. The samples were sequenced at Edinburgh Genomics using Sequel (version 2.1) chemistry.

Gene/transcript model reconstruction

Illumina raw RNA-Seq reads were quality checked and filtered using FastQC v0.11.8 (FastQC, RRID:SCR_014583) [84] and Trimmomatic v0.38 (Trimmomatic, RRID:SCR_011848) [85], respectively. Corrected reads were mapped to the genome using HISAT2 v2.2.0 (HISAT2, RRID:SCR_015530) [86, 87] and transcript models assembled using StringTie2 v2.1.1 (StringTie, RRID:SCR_0 16323) [50]. The resulting transcript models file was converted with tama_format_gtf_to_bed12_stringtie.py. Hereinafter, all tools described as starting with "tama_" are part of the software suite Transcriptome Annotation by Modular Algorithms [88], except for tama_merge_report_parser.pl [89].

PacBio raw Iso-Seq reads were pre-processed using the IsoSeq3 pipeline to obtain full-length, non-chimeric reads (FLNC; first 3 steps in [90]; ccs v3.3.0, lima v1.8.0, refine v3.1.0). Afterwards, fasta sequences were extracted from bam files using Bamtools v2.5.1 (Bamtools, RRID:SCR_015987) [91] and poly-A tails were trimmed using tama_flnc_polya_cleanup.py (v20191022). These FLNC were mapped to the reference genome with the splice site-aware mapper Minimap2 v2.17-r974-dirty. Redundant transcript models were collapsed with the capped flag (-x capped) using tama_collapse.py when coverage was ≥95% (-c 95). Additionally, 5' threshold and 3' threshold (tolerance in bp for grouping reads) were set to 100 (-a 100 -z 100).

Functional annotation

Transcript models from all 6 tissues inferred by the short-read and long-read pipelines were merged on the basis of similarity using tama_merge.py (options -a 100 -z 100 -d merge_dup) with different priorities for splice junctions and transcript end sites. Short-read inferred transcript models were given higher priority on splice junctions, whereas long-read inferred transcript models were given higher priority on transcript end sites. Nucleotide sequences based on coordinates in the merged transcriptome were extracted from the reference genome using Bedtools v2.29.0 (BEDTools, RRID:SCR_006646) [92]. The protein-coding potential was predicted with CPC2 v0.1 (Coding Potential Calculator, RRID:SCR_002764) [93] based on the transcripts' sequence features. ORFs were predicted and translated into amino acid sequences using tama_orf_seeker.py. Putative protein-coding sequences were identified in the UniProt/UniRef50 database v2019_10 (UniRef, RRID:SCR_010646) [94] using Blastp v2.9.0 (BLASTP, RRID:SCR_001010) [95]. The results were filtered for top hits with tama_orf_blastp_parser.py, and a new annotation with coding sequence (CDS) regions was created using tama_cds_regions_bed_add.py.

Identification of RIG-I/DDX58

The nucleotide sequence of the antiviral innate immune response receptor RIG-I/DDX58 in mallard (Anas platyrhynchos, version NP_001297309.1) was used to search the tufted duck genome assembly using default Tblastn (TBLASTN, RRID:SCR_011822) [96] settings on NCBI. The protein sequences of the mallard RIG-I/DDX58 were downloaded (2 isoforms) and the predicted tufted duck ORFs searched in these using Blastp (v2.10.0+). Matching ORFs were aligned with the mallard isoforms using Clustal (Clustal 2, RRID:SCR_017055) [97], and protein variation analysed using PROVEAN (PROVEAN, RRID:SCR_002182) [98] and SIFT (SIFT, RRID:SCR_012813) [99]. Transcription factor binding sites (TFBS) were identified 1 kb upstream of RIG-I/DDX58 in each species and compared by searching the TRANSFAC database (TRANS-FAC, RRID:SCR_005620) [100] with the software P-Match (Gene Regulation Programs, RRID:SCR_007787) [101].

Tissue-specific expression analysis

In addition to merging transcript models, tama_merge.py also creates gene and transcript reports that trace the source (in this case: pipeline and tissue) of each gene and transcript, respectively. The gene report was parsed with tama_merge_report_parser.pl [89] and filtered for genes found in UniRef50: Each gene was assigned a binary TRUE or FALSE label depending on the support of each of the 12 possible sources (2 pipelines and 6 tissues). The result was loaded into UpSetR [102, 103] to visualize intersections of tissue-specific evidence for an expressed gene in each pipeline.

Small RNA analyses

Illumina raw reads were quality checked with FastQC v0.11.8 and adapters removed with Cutadapt v2.10 (cutadapt, RRID: SCR_011841) [104]. Corrected reads were mapped to the reference genome using STAR v2.7.3a (STAR, RRID:SCR_004463) [105] and assembled with Cufflinks v2.2.1 (Cufflinks, RRID:SCR_01459 7) [106-109]. Nucleotide sequences were extracted from the reference genome at the assembled transcripts' coordinates using Bedtools getfasta -split (v2.29.2), and transcript lengths extracted from the output of Samtools faidx (SAMTOOLS, RRID: SCR_002105) [110]. All plots were created using the package ggplot2 (ggplot2, RRID:SCR_014601) [111] in RStudio (RStudio, RRID: SCR_000432) [103, 112].

The tool cmscan v1.1.3 from the software suite Infernal (Infernal, RRID:SCR_011809) [113] was used to predict structural RNAs in the reference genome (in silico) and to annotate assembled small RNA transcripts (in vitro) based on Rfam (Rfam, RR ID:SCR_007891) [114, 115] covariance models downloaded from

The output of cmscan (tblout) was converted to gff3 annotation files using tblout2gff3.pl [89]. Shared intervals between in silico and in vitro annotations were identified with the intersect option of Bedtools v2.29.2 [92].

Availability of Source Code and Requirements

Perl and R scripts used in this study are available on GitLab at https://gitlab.com/rcmueller/tufted_duck_annotation under MIT license.

Data Availability

The data sets supporting the results of this article are available in NCBI and Figshare. The curated assembly of the tufted duck genome has been deposited in NCBI under accession No. GCF_009819795.1 [44]. The Illumina RNA-Seg and small RNA-Seg, and PacBio Iso-Seg raw reads have been deposited in NCBI under BioProject PRJNA561952 (SRA accession Nos. SRX9968577-SRX9968594). Supporting data have been deposited on Figshare: Illumina RNA-Seq transcript models [117], PacBio Iso-Seq transcript models [118], functional annotation [119], expression atlas [120], Illumina small RNA-Seq annotation [121], additional scripts [89]. All supporting data and materials are available in the GigaScience GigaDB database [122].

Additional Files

Supplementary Figure S1. Hi-C contact map of Aythya fuliqula visualised in HiGlass.

Supplementary Figure S2. Comparison of genomic repeat content between tufted duck and the mallard.

Supplementary Figure S3. Comparison of repeat and unique sequence content in orthologous chromosome pairs between tufted duck and mallard.

Supplementary Figure S4. Identified telomeric and centromeric repeats in the tufted duck genome assembly.

Supplementary Figure S5. Distribution of single- and multi-exon transcripts per tissue and pipeline.

Supplementary Figure S6. Protein-coding potential calculated

Supplementary Figure S7. Pipeline-tissue-specific expression of genes found in UniProt.

Supplementary Figure S8. Length distribution of small RNA transcripts.

Supplementary Figure S9. Orthologous chromosome pairs of tufted duck and mallard.

Supplementary Table S1. Assembly metrics before and after cu-

Supplementary Table S2. Assembly and annotation statistics. Supplementary Table S3. Gene prediction and transcript model reconstruction.

Supplementary Table S4. Protein variation effect analysis.

Supplementary Table S5. SIFT amino acid substitution analysis. Supplementary Table S6. Transcript factor binding sites analy-

Supplementary Table S7. RNA extraction details.

Supplementary Table S8. Small RNA extraction details.

Supplementary Table S9. cDNA library details.

Supplementary Listing 1. Full alignment of antiviral immune response receptor (Aythya fuligula and Anas platyrhynchos).

Abbreviations

AIV: avian influenza A virus; BLAST: Basic Local Alignment Search Tool; bp: base pairs; cDNA: complementary DNA; CLR: continuous long reads; DLS: direct label and stain; FLNC: fulllength, non-chimeric reads; Gb: gigabase pairs; HPAIV: highly pathogenic avian influenza virus; kb: kilobase pairs; LPAIV: low pathogenic avian influenza virus; Mb: megabase pairs; NCBI: National Center for Biotechnology Information; NGS: next-generation sequencing; nt: nucleotides; ORF: open reading frame; PacBio: Pacific Biosciences; SRA: Sequence Read Archive; TAMA: Transcriptome Annotation by Modular Algorithms; VGL: Vertebrate Genomes Lab; VGP: Vertebrate Genomes Project; ZMW: zero-mode waveguide.

Ethical Approval

All animal experiments were carried out in strict accordance with a protocol legally approved by the regional board of the animal ethics committee, Sweden (permission No. 5.8.18-07998/2017). The animal experiments were conducted in biosafety level 2 animal facilities at the Swedish National Veterinary Institute (Uppsala, Sweden).

Competing Interests

The authors declare that they have no competing interests.

Funding

The authors gratefully acknowledge funding of a Short-Term Scientific Mission (STSM) through COST Action CA15112 - Functional Annotation of Animal Genomes - European network (FAANG-Europe), and funding from the Ministry of Science, Research and the Arts Baden-Württemberg (MWK) during the Science Data Centre project BioDATEN.

Authors' Contributions

R.C.M.: Conceptualization, data curation, formal analysis, investigation, methodology, project administration, software, visualization, writing-original draft, writing-review and editing; P.E.: Resources, writing-original draft; K.H.: Formal analysis, methodology, resources, validation, visualization, writingoriginal draft; M.U.S.: Formal analysis, methodology, validation, writing-original draft; R.I.K.: Conceptualization, methodology, software; K.M.: Methodology, writing-original draft; A.W.: Resources; O.F.: Formal analysis; B.H.: Formal analysis; J.M.: Formal analysis; W.C.: Formal analysis; J.T.: Formal analysis; J.M.D.W.: Formal analysis; J.D.J.: Resources; M.M.N.: Resources; B.O.: Funding acquisition; E.D.J.: Project administration, resources; J.S.: Project administration, resources, supervision; L.E.: Formal analysis, investigation, visualization; R.H.S.K.: Conceptualization, funding acquisition, project administration, resources, supervision; all authors read and approved the final manuscript.

Acknowledgements

The authors acknowledge support by the local highperformance computing resources through the core facility SCCKN at the University of Konstanz. The authors also acknowledge continued support by the Max Planck Insitute of Animal Behavior and its director Martin Wikelski.

References

- 1. Atkinson PW, Clark JA, Delany S, et al. Urgent Preliminary Assessment of Ornithological Data Relevant to the Spread of Avian Influenza in Europe. Wageningen: Wetlands International; 2006. https://library.wur.nl/WebQuery/wurpub s/369934. Accessed 17 November, 2020.
- Huang ZYX, Xu C, van Langevelde F, et al. Contrasting effects of host species and phylogenetic diversity on the occurrence of HPAI H5N1 in European wild birds. J Anim Ecol 2019;88(7):1044-53.
- Kraus RHS. The role of mallard (Anas platyrhynchos) in the spread of avian influenza: Genomics, population genetics, and flyways. Ph.D. Thesis, Wageningen University. 2011.
- Olsen B, Munster VJ, Wallensten A, et al. Global patterns of influenza A virus in wild birds. Science 2006;312(5772):384-
- Webster RG, Yakhno M, Hinshaw VS, et al. Intestinal influenza: Replication and characterization of influenza viruses in ducks. Virology 1978; 84(2):268-78.
- Prum RO, Berv JS, Dornburg A, et al. A comprehensive phylogeny of birds (Aves) using targeted next-generation DNA sequencing. Nature 2015;526(7574):569-73.
- Verhagen JH, Eriksson P, Leijten L, et al. Host range of influenza A virus H1 to H16 in Eurasian ducks based on tissue and receptor binding studies. J Virol 2021;95(6):doi:10.1128/JVI.01873-20.
- Abdo W, Haridy M, Katou Y, et al. Pathological and immunohistochemical findings of natural highly pathogenic avian influenza infection in tufted ducks during 2010-2011 outbreaks in Japan. J Vet Med Sci 2014;76(9):1285-90.
- Bröjer C, van Amerongen G, van de Bildt M, et al. Pathogenicity and tissue tropism of currently circulating highly pathogenic avian influenza A virus (H5N1; Clade 2.3.2) in tufted ducks (Aythya fuliqula). Vet Microbiol 2015;180(3):273-80.
- 10. Bröjer C, Ågren EO, Uhlhorn H, et al. Pathology of natural highly pathogenic avian influenza H5N1 infection in wild tufted ducks (Aythya fuliqula). J Vet Diagn Invest 2009;21(5):579-87.
- 11. Fiedler W, Bauer HG. "Massive" outbreak of high pathogenic avian influenza among wild ducks at Lake Constance in autumn 2016. In: 5th Pan-European Duck Symposium, Isle of Great Cumbrae, Scotland; 2018. https://ducksg.org/events/ peds5/. Accessed: 30 September, 2020.
- 12. Kraus RH, Kerstens HH, van Hooft P, et al. Widespread horizontal genomic exchange does not erode species barriers among sympatric ducks. BMC Evol Biol 2012;12:45.
- 13. Ottenburghs J, Ydenberg RC, Van Hooft P, et al. The Avian Hybrids Project: Gathering the scientific literature on avian hybridization. Ibis 2015;157(4):892-4.
- 14. Jourdain E, van Riel D, Munster VJ, et al. The pattern of influenza virus attachment varies among wild bird species. PLoS One 2011;6(9):e24155
- 15. Evseev D, Magor KE. Innate immune responses to avian influenza viruses in ducks and chickens. Vet Sci 2019;6(1):5.
- 16. Huang Y, Li Y, Burt DW, et al. The duck genome and transcriptome provide insight into an avian influenza virus reservoir species. Nat Genet 2013;45(7):776-83.

- 17. Kraus RHS, Wink M. Avian genomics: Fledging into the wild! J Ornithol 2015;156(4):851-65.
- 18. Dheilly NM, Adema C, Raftos DA, et al. No more nonmodel species: The promise of next generation sequencing for comparative immunology. Dev Comp Immunol 2014;45(1):56-66.
- 19. . In: Vignal A, Eory L. Kraus RHS. Avian genomics in animal breeding and the end of the model organism, ed. Avian Genomics in Ecology and Evolution: From the Lab into the Wild. Cham: Springer; 2019:21-67.
- Globig A, Staubach C, Beer M, et al. Epidemiological and ornithological aspects of outbreaks of highly pathogenic avian influenza virus H5N1 of Asian lineage in wild birds in Germany, 2006 and 2007. Transbound Emerg Dis 2009;56(3):57-72.
- 21. Taylor LH, Latham SM, Woolhouse ME. Risk factors for human disease emergence. Philos Trans R Soc Lond B Biol Sci 2001;356(1411):983-9.
- van Doorn HR. Emerging infectious diseases. Medicine 2017;45(12):798-801.
- 23. Ekblom R, Galindo J. Applications of next generation sequencing in molecular ecology of non-model organisms. Heredity 2011;107(1):1-15.
- Rhie A, McCarthy SA, Fedrigo O, et al. Towards complete and error-free genome assemblies of all vertebrate species. Nature 2021;592(7856):737-46.
- Jax E, Wink M, Kraus RHS. Avian transcriptomics: opportunities and challenges. J Ornithol 2018;159(3):599-629.
- Kuo RI, Tseng E, Eory L, et al. Normalized long read RNA sequencing in chicken reveals transcriptome complexity similar to human. BMC Genomics 2017;18:323.
- 27. Warren WC, Hillier LW, Tomlinson C, et al. A new chicken genome assembly provides insight into avian genome structure. G3 (Bethesda) 2017;7(1):109-17.
- 28. Steijger T, Abril JF, Engström PG, et al. Assessment of transcript reconstruction methods for RNA-Seq. Nat Methods 2013;10(12):1177-84.
- Korlach J, Gedman G, Kingan SB, et al. De novo PacBio longread and phased avian genome assemblies correct and add to reference genes generated with intermediate and short reads. Gigascience 2017;6:doi:10.1093/gigascience/gix085
- Au KF, Sebastiano V, Afshar PT, et al. Characterization of the human ESC transcriptome by hybrid sequencing. Proc Natl Acad Sci U S A 2013;110(50):E4821-30.
- 31. Sharon D, Tilgner H, Grubert F, et al. A single-molecule longread survey of the human transcriptome. Nat Biotechnol 2013;31(11):1009-14.
- 32. Li Y, Dai C, Hu C, et al. Global identification of alternative splicing via comparative analysis of SMRT- and Illuminabased RNA-Seq in strawberry. Plant J 2017;90(1):164-76.
- 33. Reixachs-Solé M, Ruiz-Orera J, Albà MM, et al. Ribosome profiling at isoform level reveals evolutionary conserved impacts of differential splicing on the proteome. Nat Commun 2020;11(1):1768.
- 34. Zhang H, Jain C, Aluru S. A comprehensive evaluation of long read error correction methods. BMC Genomics 2020;21(6):889.
- 35. Chapman JR, Hellgren O, Helin AS, et al. The evolution of innate immune genes: Purifying and balancing selection on β -defensins in waterfowl. Mol Biol Evol 2016;33(12):3075–87.
- 36. Karawita A, Cheng Y, Tong M, et al. Comparative genomics and transcriptomics help unravel why Australian black swans are uniquely susceptible to highly pathogenic avian influenza (HPAI). In: 7th ESWI Influenza Conference; 2020.

- https://www.researchgate.net/publication/346003425_C omparative_genomics_and_transcriptomics_help_unravel _why_Australian_Black_Swans_are_uniquely_susceptible_t o_highly_pathogenic_avian_influenza_HPAI, Accessed: 9 February, 2021.
- 37. Mueller RC, Mallig N, Smith J, et al. Avian Immunome DB: An example of a user-friendly interface for extracting genetic information. BMC Bioinformatics 2020;21(1):502.
- 38. Howe K, Chow W, Collins J, et al. Significantly improving the quality of genome assemblies through curation. Gigascience 2021;10:doi:10.1093/gigascience/giaa153.
- 39. Naguib MM, Eriksson P, Jax E, et al. Revealing interspecies transmission barriers of avian influenza A viruses. bioRxiv 2020:doi:10.1101/2020.11.17.386755.
- 40. Gregory TR. Animal Genome Size Database; 2021. https:// www.genomesize.com. Accessed: 14 July, 2021.
- 41. Melters DP, Bradnam KR, Young HA, et al. Comparative analysis of tandem repeats from hundreds of species reveals unique insights into centromere evolution. Genome Biol 2013;14(1):R10.
- 42. Hammar B. The karyotypes of nine birds. Hereditas 1966;55(2-3):367-85.
- 43. GenomeArk. Aythya fuliqula. 2019. https://vgp.github.io/ge nomeark/Aythya_fuligula/. Accessed: 2 February, 2021.
- 44. VGP. bAytFul2.Pri Genome Assembly NCBI; 2019. http s://www.ncbi.nlm.nih.gov/assembly/GCF_009819795.1. Accessed: 2 February, 2021.
- 45. GDV. Chr1: 1-207.0M Genome Data Viewer; 2019. https://www.ncbi.nlm.nih.gov/genome/gdv/browser/ge nome/?id=GCF_009819795.1. Accessed: 2 February, 2021.
- 46. Li J, Zhang J, Liu J, et al. A new duck genome reveals conserved and convergently evolved chromosome architectures of birds and mammals. Gigascience 2021;10:doi:10.1093/gigascience/giaa142.
- 47. ZJU1.0 Genome Assembly NCBI. 2020. https://www. ncbi.nlm.nih.gov/assembly/GCF_015476345.1. Accessed: 2 February, 2021.
- Genome Reference Consortium. GRCg6a galGal6 Genome - Assembly - NCBI. 2018.
- 49. VGP. bTaeGut2.Pat.W.v2 Genome Assembly NCBI. 2020. https://www.ncbi.nlm.nih.gov/assembly/GCF_008822105.2. Accessed: 2 February, 2021.
- 50. Kovaka S, Zimin AV, Pertea GM, et al. Transcriptome assembly from long-read RNA-Seq alignments with StringTie2. Genome Biol 2019;20(1):278.
- 51. Yin Z, Zhang F, Smith J, et al. Full-length transcriptome sequencing from multiple tissues of duck, Anas platyrhynchos. Sci Data 2019;6(1):275.
- 52. NCBI. The NCBI Eukaryotic Genome Annotation Pipeline. 2018. https://www.ncbi.nlm.nih.gov/genome/annotation_e uk/process/. Accessed: 2 February, 2021.
- 53. NCBI. Aythya Fuligula Annotation Report. 2020. https://www.ncbi.nlm.nih.gov/genome/annotation_euk/A ythya_fuligula/100/#TranscriptAlignmentStats. Accessed: 2 February, 2021.
- 54. NCBI. Anas Platyrhynchos Annotation Report. 2020. https://www.ncbi.nlm.nih.gov/genome/annotation_euk/A nas_platyrhynchos/104/. Accessed: 2 February, 2021.
- 55. NCBI. Gallus Gallus Annotation Report. 2018. https: //www.ncbi.nlm.nih.gov/genome/annotation_euk/Gallus_g allus/104/. Accessed: 2 February, 2021.
- 56. NCBI. Taeniopygia Guttata Annotation Report. 2020. https://www.ncbi.nlm.nih.gov/genome/annotation_euk/T aeniopygia_guttata/105/. Accessed: 2 February, 2021.

- 57. Kalvari I, Nawrocki EP, Ontiveros-Palacios N, et al. Rfam 14: Expanded coverage of metagenomic, viral and microRNA families. Nucleic Acids Res 2021;49(D1):D192-200.
- Bartel DP. MicroRNAs: Genomics, biogenesis, mechanism, and function. Cell 2004;116(2):281-97.
- Lee RC, Feinbaum RL, Ambros V. Cell 1993;75(5):843-54.
- 60. Long K, Feng S, Ma J, et al. Small non-coding RNA transcriptome of four high-altitude vertebrates and their lowaltitude relatives. Sci Data 2019;6(1):192.
- Baroin-Tourancheau A, Jaszczyszyn Y, Benigni X, et al. Evaluating and correcting inherent bias of microRNA expression in Illumina sequencing analysis. Front Mol Biosci 2019;6:doi:10.3389/fmolb.2019.00017.
- 62. Fuchs RT, Sun Z, Zhuang F, et al. Bias in ligation-based small RNA sequencing library construction is determined by adaptor and RNA structure. PLoS One 2015;10(5):e0126049.
- Aravin AA, Lagos-Quintana M, Yalcin A, et al. The small RNA profile during Drosophila melanogaster development. Dev Cell 2003;5(2):337-50.
- Bartel DP. MicroRNAs: Target recognition and regulatory functions. Cell 2009;136(2):215-233.
- Chen X, Ba Y, Ma L, et al. Characterization of microRNAs in serum: A novel class of biomarkers for diagnosis of cancer and other diseases. Cell Res 2008;18(10):997-1006.
- Roosbroeck KV, Pollet J, Calin GA. miRNAs and long noncoding RNAs as biomarkers in human diseases. Expert Rev Mol Diagn 2013;13(2):183-204.
- 67. Web of Science. Web of Science [v.5.35] Web of Science Core Collection Basic Search. 2021. https: //apps.webofknowledge.com/WOS_GeneralSearch_inp ut.do?product=WOS&search_mode=GeneralSearch&SID=F 5sA4vChp9eq4hS7GWi&preferencesSaved=. Accessed: 2 February, 2021.
- Fridrich A, Hazan Y, Moran Y. Too many false targets for microRNAs: Challenges and pitfalls in prediction of miRNA targets and their gene ontology in model and non-model organisms. BioEssays 2019;41(4):1800169.
- VGP. VGP-Assembly. 2021. https://github.com/VGP/vgp-ass embly. Accessed: 22 June, 2021.
- 70. Chin CS, Peluso P, Sedlazeck FJ, et al. Phased diploid genome assembly with single-molecule real-time sequencing. Nat Methods 2016;13(12):1050-4.
- 71. Guan D, McCarthy SA, Wood J, et al. Identifying and removing haplotypic duplication in primary genome assemblies. Bioinformatics 2020;36(9):2896-8.
- 72. Ghurye J, Rhie A, Walenz BP, et al. Integrating Hi-C links with assembly graphs for chromosome-scale assembly. PLoS Comput Biol 2019;15(8):e1007273.
- 73. Pacific Biosciences. PacificBiosciences/GenomicConsensus. 2020. https://github.com/PacificBiosciences/GenomicCon sensus. Accessed: 2 February, 2021.
- 74. Garrison E, Marth G. Haplotype-based variant detection from short-read sequencing(2012). arXiv:1207.3907.
- 75. Illumina. HiSeq 3000/HiSeq 4000 Systems. https: //www.illumina.com/systems/sequencing-platforms/hi seq-3000-4000.html. Accessed: 12 November, 2021.
- 76. Illumina. NovaSeq 6000 System. https://www.illumina.com /systems/sequencing-platforms/novaseq.html. Accessed: 12 November, 2021.
- 77. Smit A, Hubley R, Green P. RepeatMasker Open-4.0. 2015. http://repeatmasker.org/. Accessed: 10 November, 2020.
- 78. Hubley R, Finn RD, Clements J, et al. The Dfam Database of repetitive DNA families. Nucleic Acids Res 2016;44(D1):D81-9.

- 79. Bao W, Kojima KK, Kohany O. Repbase Update, a database of repetitive elements in eukaryotic genomes. Mob DNA 2015;6(1):11.
- 80. Smit A, Hubley R. RepeatModeler Open-1.0. 2015. http://re peatmasker.org. Accessed: 10 November 2020.
- 81. Uno Y, Nishida C, Hata A, et al. Molecular cytogenetic characterization of repetitive sequences comprising centromeric heterochromatin in three Anseriformes species. PLoS One 2019;14(3):e0214028.
- 82. Li H. Minimap2: Pairwise alignment for nucleotide sequences. Bioinformatics 2018;34(18):3094-100.
- 83. Krzywinski M, Schein J, Birol İ, et al. Circos: An information aesthetic for comparative genomics. Genome Res 2009;19(9):1639-45.
- 84. Andrews S, Krueger F, Segonds-Pichon A, et al. FastQC. 2012. http://www.bioinformatics.babraham.ac.uk/projects /fastqc. Accessed: 23 September, 2020.
- 85. Bolger AM, Lohse M, Usadel B. Trimmomatic: A flexible trimmer for Illumina sequence data. Bioinformatics 2014;30(15):2114-20.
- 86. Kim D, Paggi JM, Park C, et al. Graph-based genome alignment and genotyping with HISAT2 and HISAT-Genotype. Nat Biotechnol 2019;37(8):907-15.
- 87. Kim D, Langmead B, Salzberg SL. HISAT: A fast spliced aligner with low memory requirements. Nat Methods
- 88. Kuo RI, Cheng Y, Zhang R, et al. Illuminating the dark side of the human transcriptome with long read transcript sequencing. BMC Genomics 2020;21(1):751.
- 89. Mueller RC. Additional scripts. 2021. https://doi.org/10.608 4/m9.figshare.13853756.v1. Accessed: 10 February, 2021.
- 90. Pacific Biosciences. IsoSeq. 2020. https://github.com/Pacific Biosciences/IsoSeq. Accessed: 2 February, 2021.
- 91. Barnett DW, Garrison EK, Quinlan AR, et al. BamTools: A C++ API and toolkit for analyzing and managing BAM files. Bioinformatics 2011;27(12):1691-2.
- 92. Quinlan AR, Hall IM. BEDTools: A flexible suite of utilities for comparing genomic features. Bioinformatics 2010;26(6):841-2.
- 93. Kang YJ, Yang DC, Kong L, et al. CPC2: A fast and accurate coding potential calculator based on sequence intrinsic features. Nucleic Acids Res 2017;45(W1):W12-6.
- 94. The UniProt Consortium. UniProt: A Worldwide Hub of Protein Knowledge. Nucleic Acids Res 2019;47(D1):D506-15.
- 95. Altschul SF, Gish W, Miller W, et al. Basic Local Alignment Search Tool. J Mol Biol 1990;215(3):403-10.
- 96. Altschul SF, Madden TL, Schäffer AA, et al. Gapped BLAST and PSI-BLAST: A new generation of protein database search programs. Nucleic Acids Res 1997;25(17):3389-402.
- 97. Larkin MA, Blackshields G, Brown NP, et al. Clustal W and Clustal X Version 2.0. Bioinformatics 2007;23(21):2947-8.
- 98. Choi Y, Chan AP. PROVEAN Web Server: A tool to predict the functional effect of amino acid substitutions and indels. Bioinformatics 2015;31(16):2745-7.
- 99. Ng PC, Henikoff S. SIFT: Predicting amino acid changes that affect protein function. Nucleic Acids Res 2003;31(13):3812-
- 100. Matys V, Fricke E, Geffers R, et al. TRANSFAC: Transcriptional regulation, from patterns to profiles. Nucleic Acids Res 2003;31(1):374-8.
- 101. Chekmenev DS, Haid C, Kel AE. P-Match: Transcription factor binding site search by combining patterns and weight matrices. Nucleic Acids Res 2005;33(suppl 2):W432-7.
- 102. Conway JR, Lex A, Gehlenborg N. UpSetR: An R package for

- the visualization of intersecting sets and their properties. Bioinformatics 2017;33(18):2938-40.
- 103. R Core Team. R: A Language and Environment for Statistical Computing. 2018. https://www.R-project.org/. Accessed: 3 April, 2020.
- 104. Martin M. Cutadapt removes adapter sequences from highthroughput sequencing reads. EMBnet J 2011;17(1):10-2.
- 105. Dobin A, Davis CA, Schlesinger F, et al. STAR: Ultrafast Universal RNA-Seq Aligner. Bioinformatics 2013; 29(1):15-21.
- 106. Roberts A, Trapnell C, Donaghey J, et al. Improving RNA-Seq expression estimates by correcting for fragment bias. Genome Biol 2011;12(3):R22.
- 107. Roberts A, Pimentel H, Trapnell C, et al. Identification of novel transcripts in annotated genomes using RNA-Seq. Bioinformatics 2011;27(17):2325-9.
- 108. Trapnell C, Hendrickson DG, Sauvageau M, et al. Differential analysis of gene regulation at transcript resolution with RNA-Seq. Nat Biotechnol 2013;31(1):46-53.
- 109. Trapnell C, Williams BA, Pertea G, et al. Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. Nat Biotechnol 2010;28(5):511-15.
- 110. Li H, Handsaker B, Wysoker A, et al. The Sequence Alignment/Map Format and SAMtools. Bioinformatics 2009;25(16):2078-9.
- 111. Wickham H. Ggplot2: Elegant graphics for data analysis. New York: Springer; 2016. https://ggplot2.tidyverse.org. Accessed: 18 November, 2020.
- 112. RStudio Team. RStudio: Integrated Development Environment for R. 2016. http://www.rstudio.com/. Accessed: 3 April, 2020.
- 113. Nawrocki EP, Eddy SR. Infernal 1.1: 100-Fold faster RNA homology searches. Bioinformatics 2013;29(22):2933-5.
- 114. Kalvari I, Nawrocki EP, Argasinska J, et al. Non-coding RNA analysis using the Rfam Database. Curr Protoc Bioinformatics 2018;62(1):e51.
- 115. Kalvari I, Argasinska J, Quinones-Olvera N, et al. Rfam 13.0: Shifting to a genome-centric resource for non-coding RNA families. Nucleic Acids Res 2018;46(D1):D335-42.
- 116. European Bioinformatics Institute. /Pub/Databases/Rfam/14.2/. http://ftp.ebi.ac.uk/pub/da tabases/Rfam/14.2/. Accessed: 17 December, 2020.
- 117. Mueller RC. Illumina RNA-Seq, Transcript Models. Figshare. 2021. https://doi.org/10.6084/m9.figshare.13850360.v1. Accessed: 10 February, 2021.
- 118. Mueller RC. PacBio Iso-Seq, Transcript Models. Figshare. 2021. https://doi.org/10.6084/m9.figshare.13850840.v1. Accessed 10 February, 2021.
- 119. Mueller RC. Functional Annotation, Illumina and PacBio Transcript Models. Figshare. 2021. https://doi.org/10.6084/ m9.figshare.13851581.v1. Accessed 10 February, 2021.
- 120. Mueller RC. Expression Atlas. Figshare.. 2021. https://doi.or g/10.6084/m9.figshare.13852802.v1. Accessed 10 February,
- 121. Mueller RC. Illumina Small RNA-Seq. Figshare. 2021. ht tps://doi.org/10.6084/m9.figshare.13852256.v1. Accessed 10 February, 2021.
- 122. Mueller RC, Ellström P, Howe K, et al. Supporting data for "A high-quality genome and comparison of short versus long read transcriptome of the palaearctic duck Aythya fuligula (tufted duck)." GigaScience Database; 2021. http://dx.doi.o rg/10.5524/100940.