Ensembl gene annotation project (e!71) Gallus gallus (chicken)

This document describes the annotation process of the high-coverage Chicken assembly, described in Figure 1. The first stage is Assembly Loading where databases are prepared and the assembly loaded into the database.

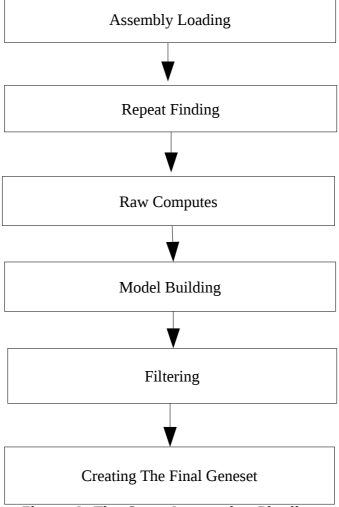


Figure 1: The Gene Annotation Pipeline

Repeat Finding

After loading into a database the genomic sequence was screened for sequence patterns including repeats using RepeatMasker [1] (version 3.2.8 with parameters '-nolow -species "gallus gallus" -s'),

RepeatModeler [2] (version open-1.0.5, to obtain a repeats library, then filtered for an additional RepeatMasker run), Dust [3] and TRF [4]. Both executions of RepeatMasker and Dust combined masked 11% of the species genome.

Raw Computes

Transcription start sites were predicted using Eponine—scan [5] and FirstEF [6]. CpG islands [Micklem, G.] longer than 400 bases and tRNAs [7] were also predicted. The results of Eponine-scan, FirstEF, CpG, and tRNAscan are for display purposes only; they are not used in the gene annotation process.

Genscan [8] was run across RepeatMasked sequence and the results were used as input for UniProt [9], UniGene [10] and Vertebrate RNA [11] alignments by WU-BLAST [12]. Passing only Genscan results to BLAST is an effective way of reducing the search space and therefore the computational resources required. This resulted in 573182 UniProt, 305991 UniGene and 296213 Vertebrate RNA sequences aligning to the genome.

cDNA and EST Alignments

Chicken cDNAs and ESTs were downloaded from ENA/Genbank/DDBJ, clipped to remove polyA tails, and aligned to the genome using Exonerate. These alignments provide supporting evidence for models.

Species	cDNA/EST		Sequences Aligned
Chicken	cDNA	51390	51368
Chicken	EST	598313	442403

Table 1: cDNA/EST alignments

All alignments were at a cut-off of 90% coverage and 80% identity.

Model Generation

Various sources of transcript and protein data were investigated and used to develop gene models using a variety of techniques. The numbers of gene models generated is described in Table 2. At this stage, many of the models may overlap. The data and techniques employed to generate models are outlined below.

Pipeline	Source	Number of Models
Targeted	5304 Uniprot chicken proteins	11511
	22457 RefSeq chicken proteins	
Similarity	193452 Uniprot PE level 1,2 proteins	209687
RNASeq	Various	7325
Ensembl Longest	16675 Ensembl Release 65 proteins for chicken,	1568
Translations	17265 Ensembl Release 65 proteins for finch,	
	14075 Ensembl Release 65 proteins for turkey	

Table 2: Gene Model Generation Overview

Targeted Pipeline: Generating coding models using species specific proteins

Protein sequences for Chicken were downloaded from public databases (UniProt SwissProt/TrEMBL [9] with Protein Existence (PE) classification level 1 or 2 and RefSeq [10]). The Chicken protein sequences were mapped to the genome using Pmatch at '-T 10'. Two sets of coding models were then produced from the proteins using Exonerate [13] and Genewise [14].

Where one protein sequence had generated more than one coding model at a locus, the BestTargeted module was used to select the coding model that most closely matched the source protein to take through to the next stage of the gene annotation process. This pipeline is shown in Figure 2.

Similarity Pipeline: Generating coding models using proteins from related species

Coding models were generated using data from related species. The UniProt alignments from the Raw Computes step were filtered and only those

sequences belonging to UniProt's Protein Existence (PE) classification level 1 and 2 were kept. WU-BLAST was rerun for these sequences and the results were passed to Genewise [14] to build coding models.

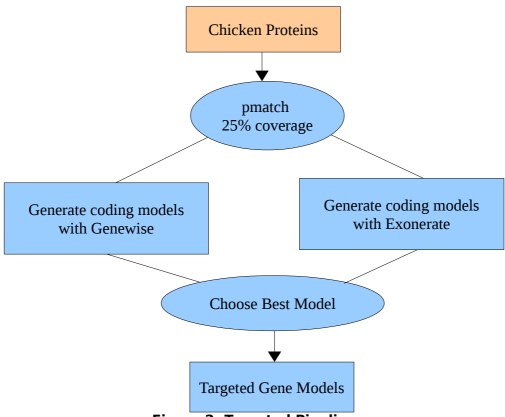


Figure 2: Targeted Pipeline

RNA-Seq Pipeline

RNA-Seq data provided by a number of sources was used in the annotation. This comprised a mixture of single and paired end data from samples including: a pool of tissues, embryo, macorphage, df1 cell lines, liver, heart, breast, cerebellum, kidney, testes and some unspecified sources. The available reads were aligned to the genome using BWA. The Ensembl RNA-Seq pipeline was used to process the BWA alignments and create further split read alignments using Exonerate.

The split reads and the processed BWA alignments were combined. The predicted open reading frames were compared to Uniprot Protein Existence (PE) classification level 1 and 2 proteins using WU-BLAST. Models with poorly scoring or no BLAST alignments were split into a seperate class.

Ensembl Longest Translations

The longest translation for each protein coding gene in Ensembl proteins release 65 for chicken, turkey and finch were downloaded. These proteins were aligned against the Chicken genome using Exonerate [13] to produce a set of coding models. The coding models were then clustered against other gene sets to filter out those overlapping gene models created from other pipelines. The remaining genes were filtered against cDNA and EST alignments to filter out those with no cDNA or EST exon support.

Filtering the Models

The filtering phase decided the subset of protein-coding transcript models, generated from the model-building pipelines, that comprise the final protein-coding gene set.

Models were filtered using the TranscriptConsensus, LayerAnnotation and GeneBuilder modules.

Apollo software [15] was used to visualise the results of filtering.

LayerAnnotation

The LayerAnnotation module was used to define a hierarchy of input data sets, from most preferred to least preferred. The output of this pipeline included all transcript models from the highest ranked input set. Models from lower ranked input sets are included only if their exons do not overlap a model from an input set higher in the hierarchy.

The top layer contained the top rated RNASeq and Targeted models generated using genewise. The remaining model sets were used in the following order:

- Similarity models built on bird proteins
- Ensembl Longest Translation models
- Other similarity models with higher level cDNA/EST support
- Targeted models generated from Exonerate
- Other similarity models with lower level cDNA/EST support
- Lower rated RNASeq models

Addition of UTR to coding models

The set of coding models not generated from RNASeq was extended into the untranslated regions (UTRs) using RNASeq, cDNA and EST sequences. At the UTR addition stage 50695 gene models out of total of 54767 non-RNASeq pipeline generated gene models had UTR added.

Generating multi-transcript genes

The above steps generated a large set of potential transcript models, many of which overlapped one another. Redundant transcript models were collapsed and the remaining unique set of transcript models were clustered into multi-transcript genes where each transcript in a gene has at least one coding exon that overlaps a coding exon from another transcript within the same gene.

At this stage the gene set comprised of 15790 genes with 16633 transcripts.

Pseudogenes

The Pseudogene module was run to identify processed pseudogenes from within the set of gene models – these were labelled as pseudogenes.

Creating The Final Gene Set

ncRNAs

Small structured non-coding genes were added using annotations taken from RFAM [16] and miRBase [17]. WU-BLAST was run for these sequences and models built using the Infernal software suite 19.

Cross-referencing

Before public release the transcripts and translations were given external references (cross-references to external databases), while translations were searched for domains/signatures of interest and labelled where appropriate. Translations were searched for signatures of interest and labelled where appropriate. Databases searched include: Seg, SignalP, Ncoils, Tmhmm, Prints, Pfscan, Pfam, Tigrfam, Superfamily, Smart and Pirsf.

Stable Identifiers

Stable identifiers were assigned to each gene, transcript, exon and translation. When annotating a species for the first time, these identifiers are auto-generated. In all subsequent annotations for a species, the stable identifiers are propagated based on comparison of the new gene set to the previous gene set. As chicken has been previously released in Ensembl a comparison was made to the previous gene set.

Final Gene Set Summary

The final gene set consists of 15508 protein coding genes, including 37 mitochondrial genes. These contain 16354 transcripts. A total of 42 pseudogenes were identified. 1558 ncRNAs were added by the ncRNA pipeline.

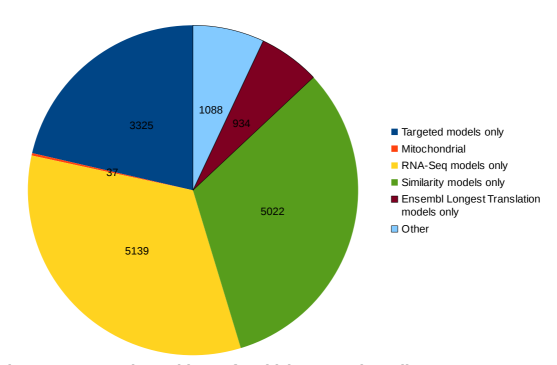


Figure 3: Supporting evidence for chicken protein coding gene set

Further information

The Ensembl gene set is generated automatically, meaning that gene models are annotated using the Ensembl gene annotation pipeline. The main focus of this pipeline is to generate a conservative set of protein-coding gene models, although non-coding genes and pseudogenes may also annotated.

Every gene model produced by the Ensembl gene annotation pipeline is supported by biological sequence evidence (see the "Supporting evidence" link on the left-hand menu of a Gene page or Transcript page); *ab initio* models are not included in our gene set. *Ab initio* predictions and the full set of cDNA and EST alignments to the genome are available on our website.

The quality of a gene set is dependent on the quality of the genome assembly. Genome assembly can be assessed in a number of ways, including:

1. Coverage estimate

- o A higher coverage usually indicates a more complete assembly.
- o Using Sanger sequencing only, a coverage of at least 2x is preferred.

2. N50 of contigs and scaffolds

- o A longer N50 usually indicates a more complete genome assembly.
- o Bearing in mind that an average human gene may be 10-15 kb in length, contigs shorter than this length will be unlikely to hold full-length gene models.

3. Number of contigs and scaffolds

- o A lower number toplevel sequences usually indicates a more complete genome assembly.
- 4. Alignment of cDNAs and ESTs to the genome

o A higher number of alignments, using stringent thresholds, usually indicates a more complete genome assembly.

More information on the Ensembl automatic gene annotation process can be found at:

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