Reviewer Report

Title: Draft genome assembly of the Bengalese finch, Lonchura striata domestica, a model for motor skill variability and learning

Version: Original Submission Date: 23 Nov 2017

Reviewer name: Morgan Wirthlin

Reviewer Comments to Author:

The study of songbirds has contributed critically to our understanding of the neural basis for learning, skilled motor behavior, sexual differentiation of the brain, and countless other topics. The recent availability of multiple high-quality avian genome assemblies has provided a starting point to explore complex questions about the genetic basis and evolution of behavior. The authors provide a high-quality genome assembly for the Bengalese finch, including a set of curated gene annotations and transcriptome data from multiple tissues. This provides a much-needed resource to the many researchers interested in this important model organism. I strongly recommend the paper for submission, following some minor revisions, which I have listed below by line number.

91 - 93 - Please split this run-on sentence into two separate sentences.

112 - "high coverage," should be high-coverage.

120 - "low coverage," should be low-coverage.

197 - Here the authors state that they manually curated models to ensure completeness and to refine UTR positions. Please provide some brief description as to the logic used to guide the curatorial process. Did the authors use aligned Cufflinks RNA-seq transcripts to guide curation, or aligned models from other species, both, neither, or something else? If it is possible to determine at this point, approximately how many gene models required manual curation?

199 - The authors state that BLASTP was used to align ORFs to the Uniprot-SwissProt protein database. Please specify the parameters used for the BLAST alignment, default or otherwise.

208 - Here the authors reference Figure 3, which provides a comparison of the Bengalese finch assembly and annotation with the assemblies from the Avian Phylogenomics Project. This figure was helpful for me to get a sense of how this assembly stacks up against previously available avian genomes. I think it would benefit the reader for the authors to provide in text some qualitative summary of the figure.

Table 1:
- Please specify the units for the Age column (e.g. post-hatch days).
- RNA libraries: the table should be alphabetized by tissue type.
- RNA libraries: one of the rows is labeled "Midbrain/brain" where it should be labeled "Midbrain/brainstem."

- RNA libraries: the NCBI records for the RNA-seq samples indicate that the muscle sample is specifically breast muscle. There's no reason not to specify "Breast muscle" in this table also.

Level of Interest

Please indicate how interesting you found the manuscript: An article of importance in its field

Quality of Written English

Please indicate the quality of language in the manuscript: Acceptable

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