

Conference Abstract

TaxonWorks as a Tool for Managing Large Biodiversity Projects

Dmitry A Dmitriev[‡], Yanghui Cao[‡], Christopher H. Dietrich[§]

[‡] University of Illinois at Urbana-Champaign, Champaign, United States of America

[§] Illinois Natural History Survey, Champaign, United States of America

Corresponding author: Dmitry A Dmitriev (arboridia@gmail.com)

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Abstract

Large systematic revisionary projects incorporating data for hundreds or thousands of taxa require an integrative approach, with a strong biodiversity-informatics core for efficient data management to facilitate research on the group. Our original biodiversity informatics platform, 3i (Internet-accessible Interactive Identification) combined a customized MS Access database backend with ASP-based web interfaces to support revisionary syntheses of several large genera of leafhoppers (Hemiptera: Auchenorrhyncha: Cicadellidae). More recently, for our National Science Foundation sponsored project, “GoLife: Collaborative Research: Integrative genealogy, ecology and phenomics of deltocephaline leafhoppers (Hemiptera: Cicadellidae), and their microbial associates”, we selected the new open-source platform [TaxonWorks](#) as the cyberinfrastructure. In the scope of the project, the original “[3i World Auchenorrhyncha Database](#)” was imported into TaxonWorks. At the present time, TaxonWorks has many tools to automatically import nomenclature, citations, and specimen based collection data. At the time of the initial migration of the 3i database, many of those tools were still under development, and complexity of the data in the database required a custom migration script, which is still probably the most efficient solution for importing datasets with long development history.

At the moment, the World Auchenorrhyncha Database comprehensively covers nomenclature of the group and includes data on 70 valid families, 6,816 valid genera, 47,064 valid species as well as synonymy and subsequent combinations (Fig. 1). In addition, many taxon records include the original citation, bibliography, type information,

are qualitative multi-state morphological descriptors) (Fig. 2). Illustrations are provided for 9,886 taxa and organized in the specialized image matrix and could be used as a pictorial key for determination of species and taxa of a higher rank.

For the phylogenetic analysis, a dataset was constructed for 730 terminal taxa and >160,000 nucleotide positions obtained using anchored hybrid enrichment of genomic DNA for a sample of leafhoppers from the subfamily Deltocephalinae and outgroups. The probe kit targets leafhopper genes, as well as some bacterial genes (endosymbionts and plant pathogens transmitted by leafhoppers). The maximum likelihood analyses of concatenated nucleotide and amino acid sequences as well as coalescent gene tree analysis yielded well-resolved phylogenetic trees (Cao et al. 2022). Raw sequence data have been uploaded to the Sequence Read Archive on [GenBank](#). Occurrence and morphological data, as well as diagnostic images, for voucher specimens have been incorporated into TaxonWorks.

Data in TaxonWorks could be exported in raw format, get accessed via Application Programming Interface (API), or be shared with external data aggregators like [Catalogue of Life](#), [GBIF](#), [iDigBio](#).

Keywords

Hemiptera, Homoptera, Auchenorrhyncha, Cicadellidae, Deltocephalinae, leafhoppers, databases, checklist, taxonomy, systematics

Presenting author

Dmitry A Dmitriev

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