

Towards unified formats for sequences, alignments, features, and annotations

Matúš Kalaš¹, Sveinung Gundersen², László Kaján³, Hervé Ménager⁴, Jon Ison⁵, Christophe Blanchet⁶, Steve Pettifer⁷, Rodrigo Lopez⁸, Kristoffer Rapacki⁵, Inge Jonassen¹, and open for contributions

¹Computational Biology Unit, Department of Informatics, University of Bergen, Bergen, Norway; ²Institute for Cancer Research, Oslo University Hospital and Department of Informatics, University of Oslo, Oslo, Norway; ³unaffiliated, previously Bioinformatics and Computational Biology Department, Technische Universität München, Garching, Germany; ⁴Institut Pasteur, Paris, France; ⁵Department of Systems Biology, Technical University of Denmark, Kongens Lyngby, Denmark; ⁶L'Institut Français de Bioinformatique, Gif-sur-Yvette, France; ⁷School of Computer Science, University of Manchester, UK; ⁸European Bioinformatics Institute, EMBL, Wellcome Trust Genome Campus, Hinxton, Cambridge, UK.



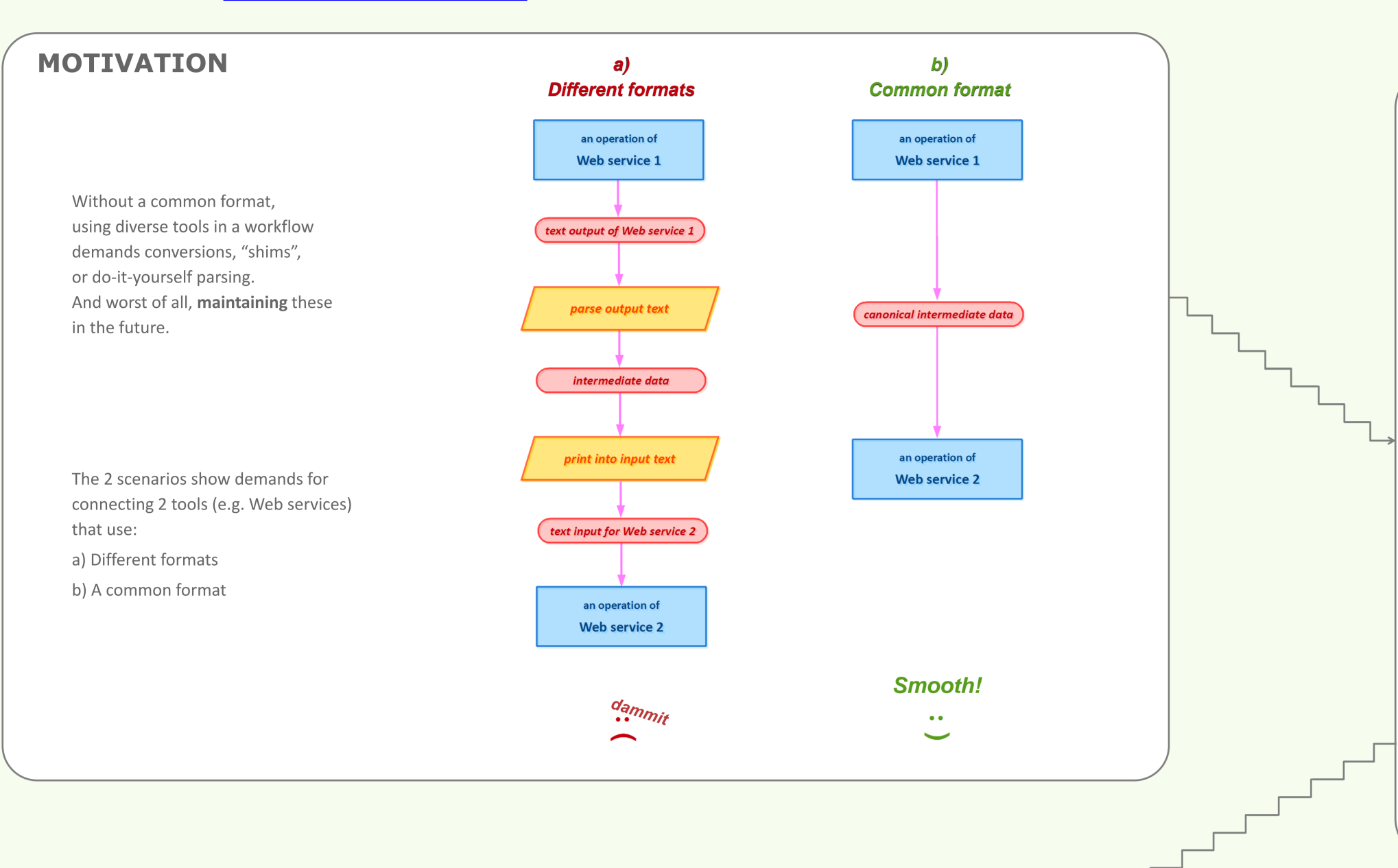


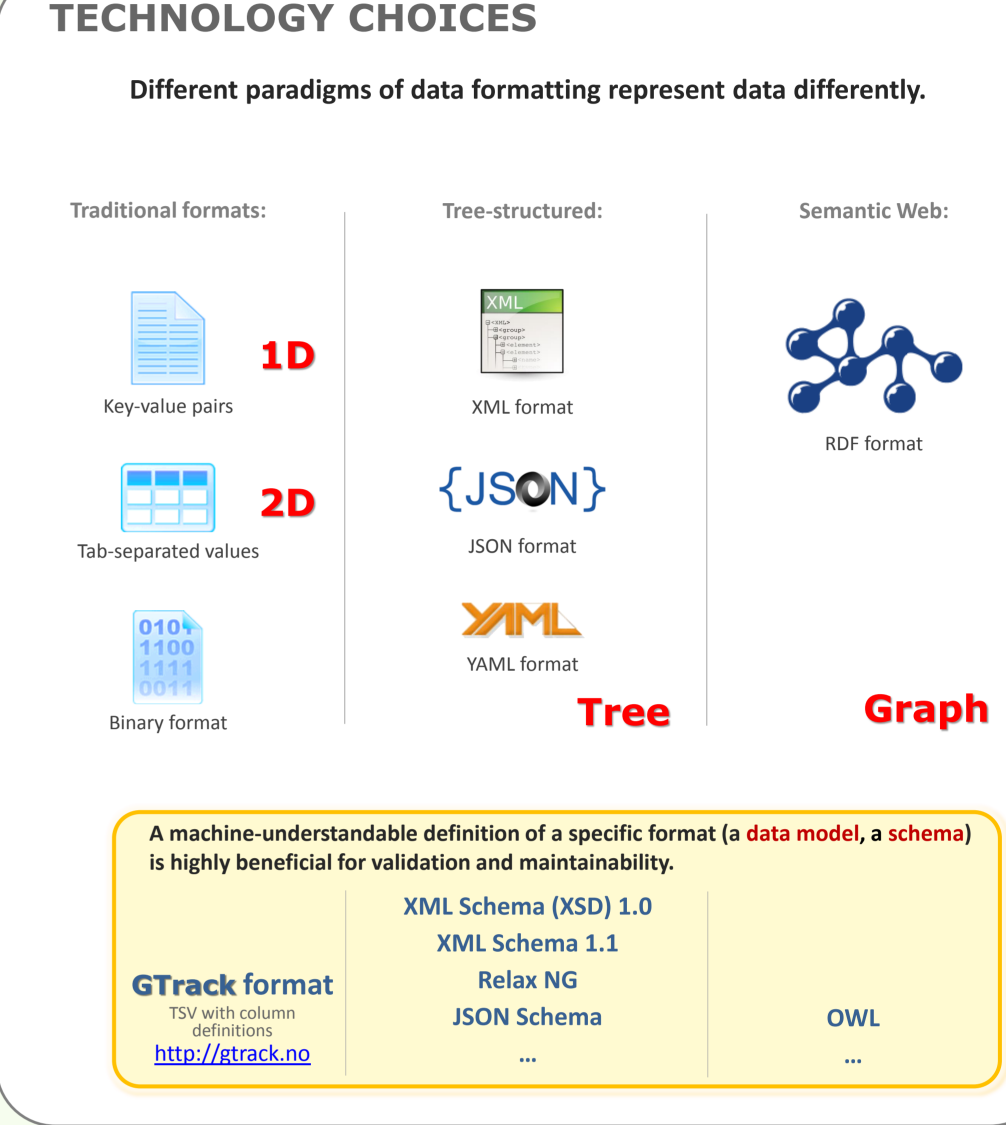
http://groups.google.com/group/bioxsd

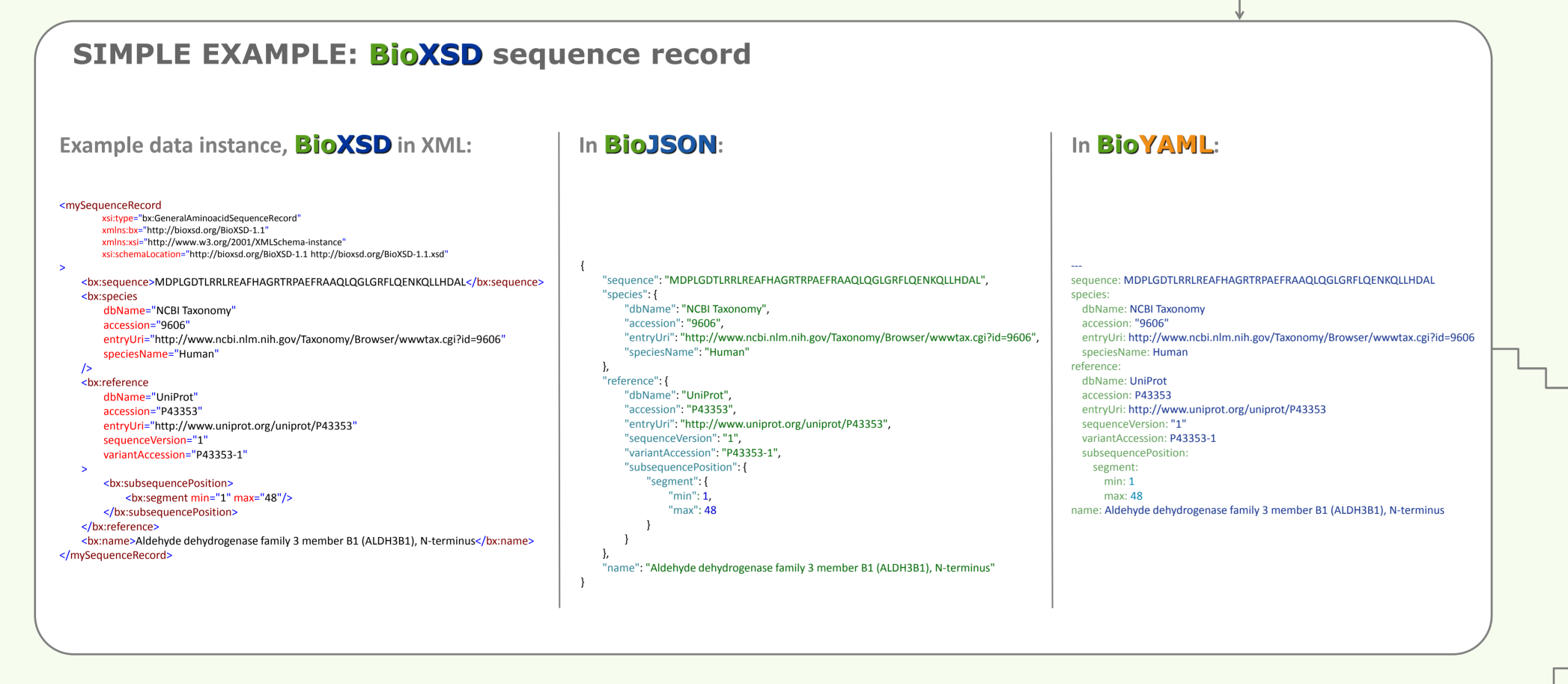


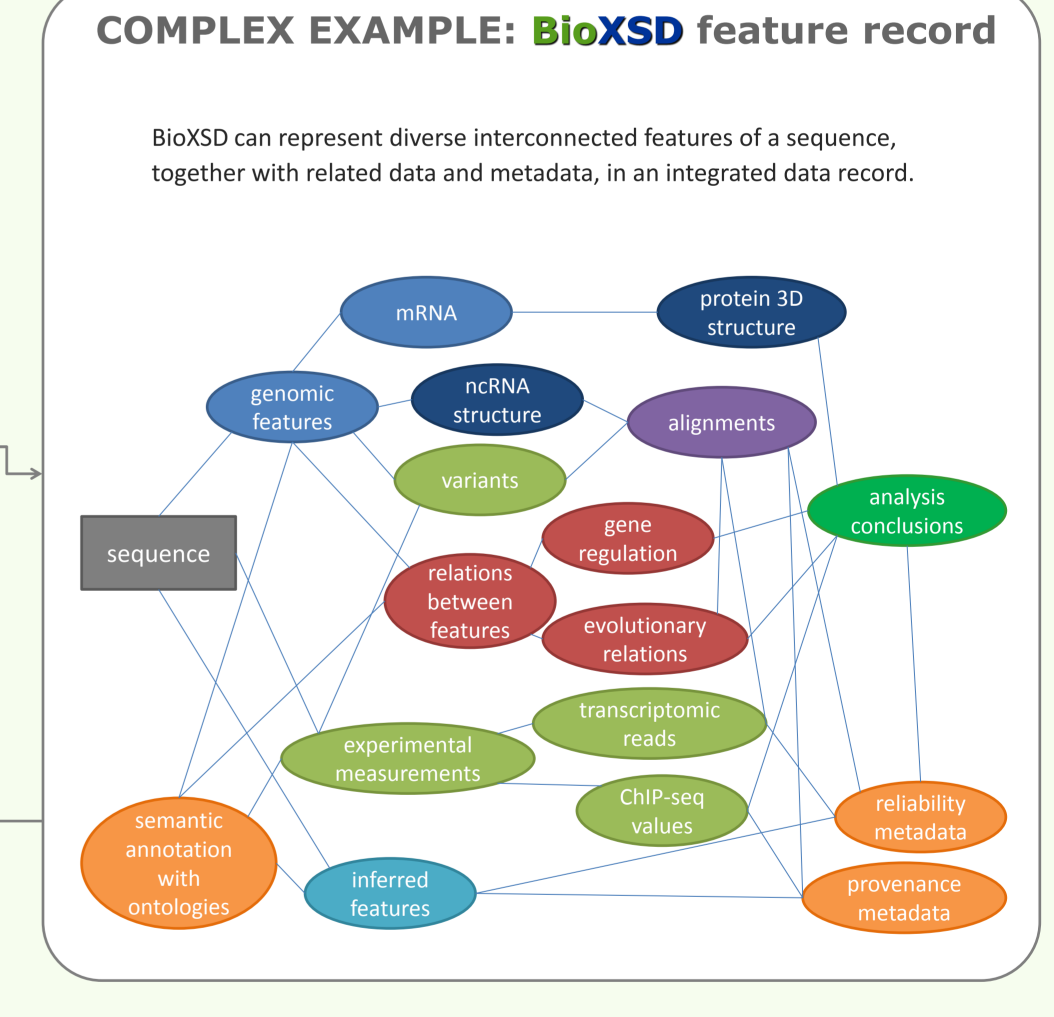
support@bioxsd.org

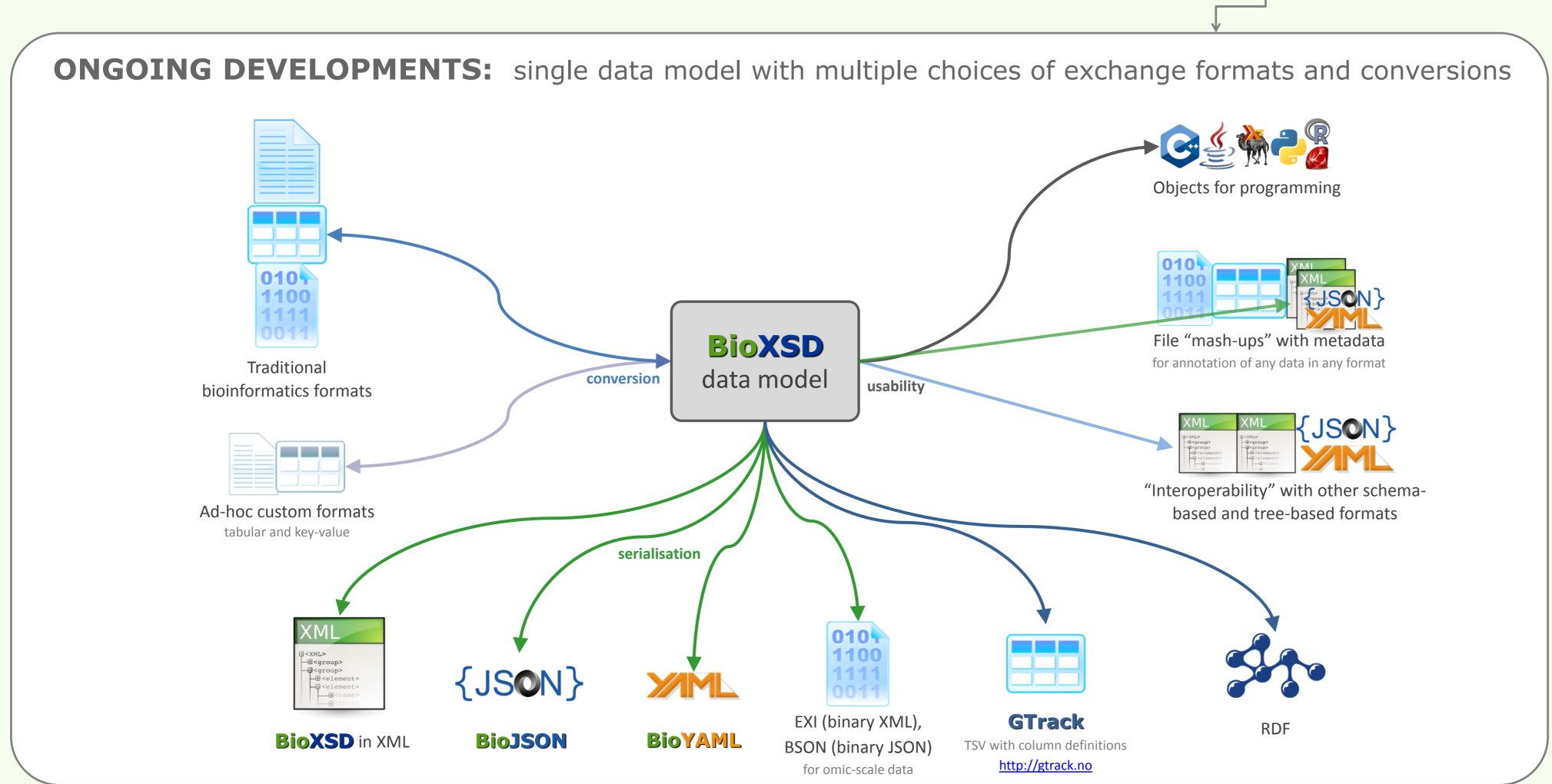
Latest stable release: http://bioxsd.org/BioXSD-1.1.xsd











BioXSD has been developed as a tree-based data model and an exchange format for basic bioinformatics data, centred around a bio-polymer sequence. BioXSD allows integration of diverse features, information, measurements, and inferred values about a biological molecule or its part, annotated with provenance and reliability metadata, ontology concepts, scientific remarks, and conclusions.

BioJSON and BioYAML are the ongoing developments. These exchange formats are based on the same data model as BioXSD, but providing serialisations in JSON and YAML respectively. BioJSON and BioYAML thus enrich the BioXSD family with alternatives to the original XML.

As tree-based data formats, BioXSD, BioJSON, and BioYAML are particularly suitable for programming in object-oriented languages, and for use with web applications and web APIs (Web services), while at the same time allowing a reasonable level of human readability.

BioXSD|BioJSON|BioYAML are developed together with GTrack (the universal tabular format for sequence features). The BioXSD|GTrack family, as a "service" developed by ELIXIR Norway, is going to support smooth interoperability between these alternative, universal formats, and between the tools that consume or provide them as inputs or outputs.











