

A reusable tree-based web-visualization to browse EDAM ontology, and contribute to it.

Bryan Brancotte¹, Christophe Blanchet², and Hervé Ménager¹

1 Bioinformatics and Biostatistics HUB, Centre de Bioinformatique, Biostatistique et Biologie Intégrative (C3BI, USR 3756 Institut Pasteur et CNRS), Paris, France 2 French Institute of Bioinformatics, CNRS IFB-Core, Gif-sur-Yvette, France

DOI: 10.21105/joss.00698

Software

■ Review 🗗

■ Repository 🗗

■ Archive ♂

Submitted: 20 February 2018 **Published:** 15 June 2018

Licence

Authors of papers retain copyright and release the work under a Creative Commons Attribution 4.0 International License (CC-BY).

Summary

The EDAM Browser, a client-side web-based visualization javascript widget that provides an interface to navigate the EDAM ontology (Ison et al. 2013). EDAM is a structured and controlled vocabulary describing bioinformatics and computational biology concepts. It is widely used to annotate resource descriptions (tools, web services) for discovery (Ison et al. 2015), workflow construction and integration. The EDAM Browser is not designed to be a generic ontology navigation and/or edition platform, a goal already achieved by many other systems such as AberOWL (Hoehndorf et al. 2015), BioPortal (Whetzel et al. 2011), OLS - Ontology Lookup Service (Jupp et al. 2015), Ontobee (Xiang et al. 2011) and WebProtégé (Tudorache et al. 2013).

Rather, the EDAM Browser is tailored to the structure and properties of EDAM, and targerts both contributors and users who might not be ontology experts. It intend to help describing resources, and to facilitate and foster community contributions to EDAM.

Availability and re-usablility

The EDAM browser is available publicly and anonymously at https://ifb-elixirfr.github.io/edam-browser/. In addition to this, its lightweight architecture makes it easy to download and run on any server or personal computer, either as a local HTML file or on a web server. It is possible to integrate the EDAM Browser and its tree representation in external websites and applications, providing a simple way for third party websites to promote EDAM-labeled resources. Both the autocomplete input field and the tree visualization are re-usable: a demonstration code is available here, showing how the tree can be integrated, how the user can interact with the tree, and how to programmaticaly interact with the tree in JavaScript.

Information display

As much as possible, the user interface aims at simplicity and relevance to the specific domain of EDAM. The creation of an interface that displays all of the information necessary to users, and avoids the use of ontology development jargon is a major goal of this project.

The interface permits the navigation between different axes of the ontology, based on the EDAM properties that define their relationships (e.g. this *format* represents this type of *data*, this *data* is an output of this *operation* or is specific of this *topic*).



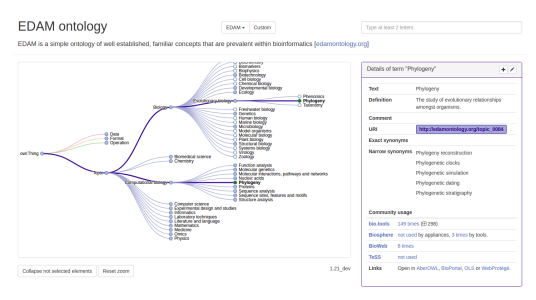


Figure 1: The term Phylogeny has two parents

We also take into account the specificities of the structure of EDAM: while being represented as a tree, it is in fact a directed acyclic graph, meaning that a term can have more than one parent. In order to improve readability when a term is selected (1) all the term's positions are shown; and (2) all paths from the root node are highlighted. A good example of this display is the Phylogeny topic) (cf Fig. 1).

On the right part of Figure 1, details of the selected concept are displayed. One last salient feature is the presentation of the usage of the concept in annotated resource collections, such as bio.tools, BioSphere (Brancotte et al. 2017), BioWeb and TeSS.

Performance and flexibility

One of the specificities of EDAM is its relatively small size in comparison with large ontologies like Gene Ontology. This reduced size makes it easy to load entirely the contents to be displayed in the browser's memory, and enables a very fast navigation, with no need to rely on server calls during this navigation (except for displaying usage statistics from external annotated resources). Using the EDAM Browser to explore a local or indevelopment version is possible. The loaded file should be formatted as a JSON file following the schema accessible here. The edam2json utility can be used to generate the ontology in this format from any EDAM owl file.

An ontology is loaded into the EDAM Browser by clicking on the button labelled *Custom* at the top of its interface, and specifying either a public URL to the file or a local path to load it from (cf Fig. 2).

Facilitating community feedback.

EDAM users can contribute their own expertise to the ontology. In order to make these contributions easier, we have implemented a form that facilitates this communication. These suggestions are automatically formatted as github issues ready to be submitted by the user (cf Fig. 3).





Figure 2: Providing a custom ontology

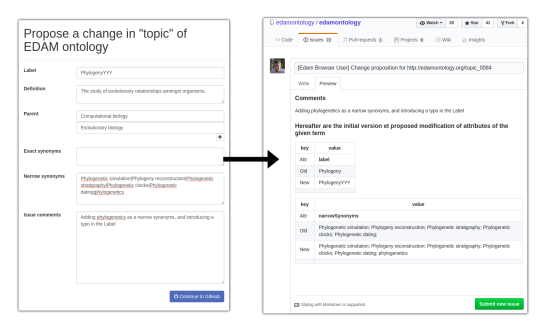


Figure 3: The edition form and the issue created



References

Brancotte, Bryan, Mohamed Bedri, Jonathan Lorenzo, Perrin Sandrine, Frédéric Séné, Awa Sepou Ngaïlo, Christophe Blanchet, and Jean-François Gibrat. 2017. "Biosphere Web: un portail haut niveau pour une utilisation bioinformatique des clouds." JOBIM (Journées Ouvertes Biologie Informatique Mathématiques). https://www.youtube.com/watch?v=gaopunfIqnc.

Hoehndorf, Robert, Luke Slater, Paul N. Schofield, and Georgios V. Gkoutos. 2015. "Aber-Owl: A Framework for Ontology-Based Data Access in Biology." *BMC Bioinformatics* 16 (1):26. https://doi.org/10.1186/s12859-015-0456-9.

Ison, Jon, Matúš Kalaš, Inge Jonassen, Dan Bolser, Mahmut Uludag, Hamish McWilliam, James Malone, Rodrigo Lopez, Steve Pettifer, and Peter Rice. 2013. "EDAM: An Ontology of Bioinformatics Operations, Types of Data and Identifiers, Topics and Formats." *Bioinformatics* 29 (10). Oxford University Press:1325–32. https://doi.org/10.1093/bioinformatics/btt113.

Ison, Jon, Kristoffer Rapacki, Hervé Ménager, Matúš Kalaš, Emil Rydza, Piotr Chmura, Christian Anthon, et al. 2015. "Tools and Data Services Registry: A Community Effort to Document Bioinformatics Resources." *Nucleic Acids Research* 44 (D1). Oxford University Press:D38–D47. https://doi.org/10.1093/nar/gkv1116.

Jupp, Simon, Tony Burdett, Catherine Leroy, and Helen E Parkinson. 2015. "A New Ontology Lookup Service at Embl-Ebi." In SWAT4LS, 118–19. {http://ceur-ws.org/Vol-1546/paper_29.pdf}.

Tudorache, Tania, Csongor Nyulas, Natalya F Noy, and Mark A Musen. 2013. "WebProtégé: A Collaborative Ontology Editor and Knowledge Acquisition Tool for the Web." Semantic Web 4 (1). IOS Press:89–99. https://doi.org/10.3233/SW-2012-0057.

Whetzel, Patricia L, Natalya F Noy, Nigam H Shah, Paul R Alexander, Csongor Nyulas, Tania Tudorache, and Mark A Musen. 2011. "BioPortal: Enhanced Functionality via New Web Services from the National Center for Biomedical Ontology to Access and Use Ontologies in Software Applications." *Nucleic Acids Research* 39 (suppl_2). Oxford University Press:W541–W545. https://doi.org/10.1093/nar/gkr469.

Xiang, Zuoshuang, Chris Mungall, Alan Ruttenberg, and Yongqun He. 2011. "Ontobee: A Linked Data Server and Browser for Ontology Terms." In *ICBO*. https://doi.org/10.1093/nar/gkw918.