



University of Dundee

# **MSAViewer**

Yachdav, Guy; Wilzbach, Sebastian; Rauscher, Benedikt; Sheridan, Robert; Sillitoe, Ian; Procter, James; Lewis, Suzanna E.; Rost, Burkhard; Goldberg, Tatyana

Published in: **Bioinformatics** 

DOI: 10.1093/bioinformatics/btw474

Publication date: 2016

**Document Version** Publisher's PDF, also known as Version of record

Link to publication in Discovery Research Portal

Citation for published version (APA):

Yachdav, G., Wilzbach, S., Rauscher, B., Sheridan, R., Sillitoe, I., Procter, J., ... Goldberg, T. (2016). MSAViewer: interactive JavaScript visualization of multiple sequence alignments. Bioinformatics. DOI: 10.1093/bioinformatics/btw474

#### **General rights**

Copyright and moral rights for the publications made accessible in Discovery Research Portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

Users may download and print one copy of any publication from Discovery Research Portal for the purpose of private study or research.
You may not further distribute the material or use it for any profit-making activity or commercial gain.
You may freely distribute the URL identifying the publication in the public portal.

#### Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Bioinformatics, 2016, 1–3 doi: 10.1093/bioinformatics/btw474 Advance Access Publication Date: 13 July 2016 Applications Note

## OXFORD

# Sequence analysis

# MSAViewer: interactive JavaScript visualization of multiple sequence alignments

Guy Yachdav,<sup>1,2</sup> Sebastian Wilzbach,<sup>1</sup> Benedikt Rauscher,<sup>1</sup> Robert Sheridan,<sup>3</sup> Ian Sillitoe,<sup>4</sup> James Procter,<sup>5</sup> Suzanna E. Lewis,<sup>6</sup> Burkhard Rost<sup>1,2</sup> and Tatyana Goldberg<sup>1,\*</sup>

<sup>1</sup>Bioinformatik - I12, TUM, Garching, 85748, Germany, <sup>2</sup>Biosof LLC, New York, NY 10001, USA, <sup>3</sup>Department of Systems Biology, Harvard Medical School, Boston, MA 02115, USA, <sup>4</sup>Institute of Structure and Molecular Biology, University College London, London, UK, <sup>5</sup>Biological Chemistry and Drug Discovery, University of Dundee, Dundee, UK and <sup>6</sup>Lawrence Berkeley National Laboratory, Berkeley, USA

\*To whom correspondence should be addressed. Associate Editor: John Hancock

Received on April 16, 2016; revised on June 3, 2016; accepted on June 29, 2016

### Abstract

**Summary:** The MSAViewer is a quick and easy visualization and analysis JavaScript component for Multiple Sequence Alignment data of any size. Core features include interactive navigation through the alignment, application of popular color schemes, sorting, selecting and filtering. The MSAViewer is 'web ready': written entirely in JavaScript, compatible with modern web browsers and does not require any specialized software. The MSAViewer is part of the BioJS collection of components.

Availability and Implementation: The MSAViewer is released as open source software under the Boost Software License 1.0. Documentation, source code and the viewer are available at http://msa.biojs.net/.

**Supplementary information:** Supplementary data are available at *Bioinformatics* online. **Contact:** msa@bio.sh

### **1** Introduction

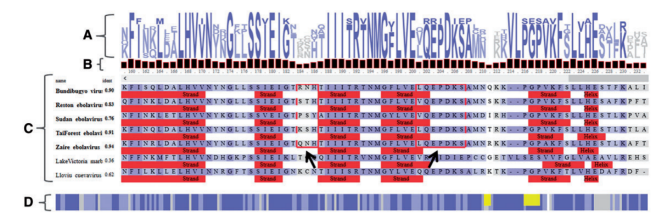
Multiple Sequence Alignment (MSA) is a fundamental procedure to capture similarities between sequences of nucleotides (DNA/RNA) or of amino acids (protein). Biologically meaningful MSAs highlight and capture sites with significant evolutionary conservation. MSAs are essential to predict aspects of protein structure (e.g. secondary structure (Rost, 2001) or protein disorder (Schlessinger *et al.*, 2011)) and function [e.g. binding sites (Ofran *et al.*, 2007) or localization (Goldberg *et al.*, 2014)]. MSAs can be used to understand genomic rearrangements (Darling *et al.*, 2004), to derive sequence homology (Altschul *et al.*, 1990) and to identify evolutionary rates (Pupko *et al.*, 2002).

MSAs are widely used to display complex annotations relating to structure and function, and to transfer those annotations to sequences that lack annotations (Waterhouse *et al.*, 2009). Many tools are available to view and analyze MSAs, including standalone applications (Larsson, 2014; Waterhouse *et al.*, 2009) and web applets (Waterhouse *et al.*, 2009). With the recent widespread adoption of JavaScript as the leading programming language for interactive web applications, new MSA viewing tools compatible with modern web browsers have been developed and made available (Martin, 2014). BioJS is one particular collection of JavaScript components with growing applications in biology (Corpas *et al.*, 2014); it is interoperable with many other data visualization tools.

Here, we describe the BioJS MSAViewer. It is readily loaded into web pages to visualize and analyze MSA datasets of arbitrary sizes. MSAViewer implements most features commonly available in other popular MSA viewing software, including scrolling, selecting, highlighting, cross-referencing with protein feature annotations and phylogenetic trees (a detailed comparison of features is in Supplementary Table S1).

<sup>©</sup> The Author 2016. Published by Oxford University Press.

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com



**Fig. 1.** A simplified view of the MSAViewer for the sequence alignment of protein VP24 within seven viruses of *Filoviridae* family. (A) Sequence logo (Schneider and Stephens, 1990) representation with conservation patterns at each position in the MSA. (B) Bar chart showing amino acid conservation per position. (C) Main MSA panel with residues in the alignment colored according to the Percentage Identity coloring scheme (Waterhouse *et al.*, 2009). Percentage sequence identities relative to consensus (consensus sequence not shown) are listed for each sequence. Filled red rectangles indicate sequence annotations provided by the user [here: secondary structure predictions of PredictProtein (Yachdav *et al.*, 2014)]. Red frames (indicated by two black arrows) are clusters of residues of ebolavirus VP24 responsible for binding to karyopherin alpha nuclear transporters to suppress antiviral defense mechanism in human (Xu *et al.*, 2014). (D) A compact overview MSA showing a bird's eye view of the full alignment. Yellow rectangles are two highlighted clusters in the main MSA panel. Despite the overall sequence homology, the highlighted regions in ebolavirus (EBOV) proteins binding karyoprotein differ from those in Llovu cuevavirus (LLOV) and Lake Victoria marburgvirus (MARV). MARV is known to not suppress host's antiviral defense mechanism (Xu *et al.*, 2014). Though no experimental information on binding of LLOV VP24 to karyoproteins is available to date, the comparison of its sequence and structural features suggests a mechanism that is more similar to EBOV than to MARV

#### 2 Visualization

The MSAViewer loads MSA data in FASTA (Pearson, 2000) or CLUSTAL (Larkin *et al.*, 2007) formats from a user's local computer or a web server. It then draws two main Canvas panels—the main panel and the overview MSA panel (Fig. 1). The choice to use Canvas over other rendering technologies is discussed in Supplementary Material Section S1.

Navigation through the alignment is enabled through various controls. First, users can scroll or use the 'jump to a column' menu item to navigate to a certain column number. Second, users can pan within the main panel to scroll through the alignment; this has proven to be a useful feature for large alignments. Finally, a second panel—the overview panel (drawn under the main panel)—provides a 'bird's eye view' perspective over the entire alignment and can also be used for navigation.

The alignment is sortable by unique identifiers, sequence labels and sequences. The percentage of gaps and the sequence identity to the consensus sequence, which is calculated from most frequent bases at each position in the alignment, can also be used for sorting. Users can select sequences, columns, or arbitrary regions for analysis, and hide them from the main panel based on their selection, conservation to the consensus sequence or the percentage of gaps. An alignment can also be searched for motifs using a regular expression (e.g. K(K|R)RK for a nuclear localization signal), which are highlighted with a red frame if matched. Sequence position annotations, such as binding sites, are provided by a user and displayed as filled rectangles below the corresponding sequence in the alignment. Users can switch between 15 predefined color schemes. The MSAViewer exports alignments and annotations as ASCII files, and the visual representation as a publication-quality figure.

#### 3 Summary

MSAViewer is a lightweight viewer for MSAs of arbitrary size. Being JavaScript-based, it can be used on any modern web browser without installing any specialized software or add-ons. As part of BioJS, the MSAviewer can interoperate with a growing set of biological data viewers. The phylogenetic tree and the sequence logo viewers are already integrated in the current release. Integration with new MSA visualization techniques, e.g. sequence bundles (Kultys *et al.*, 2014) is planned. The MSAViewer has already been found useful and became part of Galaxy (Giardine *et al.*, 2005) https://cpt.tamu.edu/clustalw-msa-and-visualisations and JalView (Waterhouse *et al.*, 2009) http:// www.jalview.org/help/html/features/biojsmsa.html.

## Funding

This work was supported by a grant from the German Federal Ministry for Education and Research (BMBF), Ernst Ludwig Ehrlich Studienwerk and the Google Summer of Code program sponsored by Google Inc.

Conflict of Interest: none declared.

#### References

- Altschul,S.F. et al. (1990) Basic local alignment search tool. J. Mol. Biol., 215, 403–410.
- Corpas, M. et al. (2014) BioJS: an open source standard for biological visualisation - its status in 2014. F1000Res., 3, 55.
- Darling, A.C. et al. (2004) Mauve: multiple alignment of conserved genomic sequence with rearrangements. Genome Res., 14, 1394–1403.
- Giardine, B. et al. (2005) Galaxy: a platform for interactive large-scale genome analysis. Genome Res., 15, 1451–1455.
- Goldberg, T. et al. (2014) LocTree3 prediction of localization. Nucleic Acids Res., 42, W350–W355.
- Kultys, M. et al. (2014) Sequence Bundles: a novel method for visualising, discovering and exploring sequence motifs. BMC Proc., 8, S8.
- Larkin, M.A. et al. (2007) Clustal W and Clustal X version 2.0. Bioinformatics, 23, 2947–2948.
- Larsson, A. (2014) AliView: a fast and lightweight alignment viewer and editor for large datasets. *Bioinformatics*, **30**, 3276–3278.
- Martin, A.C. (2014) Viewing multiple sequence alignments with the JavaScript Sequence Alignment Viewer (JSAV). F1000Res., 3, 249.
- Ofran, Y. et al. (2007) Prediction of DNA-binding residues from sequence. Bioinformatics, 23, i347-i353.

- Pearson, W.R. (2000) Flexible sequence similarity searching with the FASTA3 program package. *Methods Mol. Biol*, **132**, 185–219.
- Pupko, T. et al. (2002) Rate4Site: an algorithmic tool for the identification of functional regions in proteins by surface mapping of evolutionary determinants within their homologues. *Bioinformatics*, 18, S71–S77.
- Rost, B. (2001) Review: protein secondary structure prediction continues to rise. J. Struct. Biol., 134, 204–218.
- Schlessinger, A. et al. (2011) Protein disorder–a breakthrough invention of evolution? Curr. Opin. Struct. Biol., 21, 412–418.
- Schneider, T.D. and Stephens, R.M. (1990) Sequence logos: a new way to display consensus sequences. Nucleic Acids Res., 18, 6097–6100.
- Waterhouse, A.M. *et al.* (2009) Jalview Version 2–a multiple sequence alignment editor and analysis workbench. *Bioinformatics*, **25**, 1189–1191.
- Xu,W. *et al.* (2014) Ebola virus VP24 targets a unique NLS binding site on karyopherin alpha 5 to selectively compete with nuclear import of phosphorylated STAT1. *Cell Host Microbe*, **16**, 187–200.
- Yachdav, G. et al. (2014) PredictProtein–an open resource for online prediction of protein structural and functional features. Nucleic Acids Res., 42, W337–W343.