

Resource Summary Report

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FaBox

RRID:SCR_005350

Type: Tool

Proper Citation

FaBox (RRID:SCR_005350)

Resource Information

URL: <http://users-birc.au.dk/biopv/php/fabox/>

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Description: Tools for splitting, joining and otherwise manipulating FASTA format sequence files. The first tools in the toolbox is for manipulating fasta headers, cropping alignments and doing some sequence comparison allowing users to combine the description of data (often in excel spreadsheets) with the actual data (often DNA sequences). Also, producing correct input files for a range of programs seems to be problematic for the average user. Hence, some converters in some of the services have been included as well as some stand-alone converters. The converters are not necessarily meant to provide the final input file, but you'll get a valid input file for Arlequin, MrBayes etc. - that you may further edit so it suit your needs. This means that you may need to combine several of the tools to finish your handling - but it keeps it relatively simple to use. Please note that FaBox is written in PHP and ONLY RUNS ON A WEBSERVER.

Abbreviations: FaBox

Synonyms: FaBox - an online fasta sequence toolbox

Resource Type: software resource

Keywords: fasta, dna, protein sequence, dna sequence, protein, sequence, php

Availability: Acknowledgement requested

Resource Name: FaBox

Resource ID: SCR_005350

Alternate IDs: OMICS_01165

Record Creation Time: 20220129T080229+0000

Record Last Update: 20260808T185833+0000

Ratings and Alerts

No rating or validation information has been found for FaBox.

No alerts have been found for FaBox.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 120 mentions in open access literature.

Listed below are recent publications. The full list is available at [NIF](#) .

Chai MJ, et al. (2026) Data collected in a citizen scientist study uncover a new species record of Phoxinus minnow for Austria. Environmental monitoring and assessment, 198(4).

Casabella-Herrero G, et al. (2026) Host preference and specialization in the genus Aphanomyces (Oomycetes) from molecular and interaction network insights. Scientific reports, 16(1).

Derouiche L, et al. (2026) Genetic evidence for the rediscovery in the wild of the critically endangered Sahara killifish Apricaphanius saourensis (Cyprinodontiformes: Aphaniidae). Die Naturwissenschaften, 113(1), 21.

Beyer R, et al. (2026) Distinct properties of human pathogenic Candida species revealed by systematic comparative phenotypic screening of clinical isolates. mSystems, 11(1), e0078625.

Kniha E, et al. (2025) Combining phylogeography and climate models to track the diversification and spread of Phlebotomus simici. Scientific reports, 15(1), 10188.

Komal M, et al. (2025) Multi-marker analysis of Fasciola gigantica from cattle and buffalo across Pakistan reveals high levels of genetic diversity and novel haplotypes. Parasitology, 152(10), 1047.

Gasulla F, et al. (2025) Algae and the city: the genetic and ecophysiological diversity of photobionts in two areas of Madrid (Spain) with contrasting levels of nitrogen pollution. Environmental science and pollution research international, 32(30), 17978.

Bernard AM, et al. (2025) Connections Across Open Water: A Bi-Organelle, Genomics-Scale Assessment of Atlantic-Wide Population Dynamics in a Pelagic, Endangered Apex Predator Shark (Isurus oxyrinchus). Evolutionary applications, 18(1), e70071.

Coello Peralta RD, et al. (2025) Identification of Ancylostoma caninum in Domestic Dogs from Ecuador via Various Techniques. Medical science monitor : international medical journal of experimental and clinical research, 31, e947069.

- Kos A, et al. (2025) Disentangling the cave centipede *Lithobius stygius* species complex through molecular phylogenetics and redescription of *L. stygius* s. str. *Scientific reports*, 15(1), 42687.
- Pran?l J, et al. (2025) Identifying the richness and evolutionary relationships of *Ranunculus* sect. *Batrachium* in its diversity centre in south-western Europe. *Scientific reports*, 15(1), 14008.
- Prakas P, et al. (2024) Low Genetic Variability of the Tundra Vole in Lithuania. *Animals : an open access journal from MDPI*, 14(2).
- Hilário S, et al. (2024) Comparative genomics reveals insights into the potential of *Lysinibacillus irui* as a plant growth promoter. *Applied microbiology and biotechnology*, 108(1), 370.
- Nehemia A, et al. (2024) Anthropogenic activities along the Lake Nyasa catchments alter the habitat and genetic diversity of a Lake Salmon, *Opsaridium microlepis*. *Heliyon*, 10(20), e39051.
- Cunha-Ferreira IC, et al. (2024) Impact of *Paenibacillus elgii* supernatant on screening bacterial strains with potential for biotechnological applications. *Engineering microbiology*, 4(3), 100163.
- Takamura K, et al. (2024) Common Food Sources for Macrobenthic Communities in Different Bottom Covers: Submerged Macrophytes and Benthic Cyanobacteria in a Japanese Temperate Lake. *Ecology and evolution*, 14(11), e70359.
- Prakas P, et al. (2024) First report of *Sarcocystis halioti* (Apicomplexa) in bearded vulture (*Gypaetus barbatus*). *Veterinary research communications*, 48(1), 541.
- Huervana JJC, et al. (2023) Preliminary assessment of genetic variation in the Japanese endemic freshwater crab, *Geothelphusa dehaani*, based on mitochondrial DNA sequences. *Biodiversity data journal*, 11, e97438.
- Liu H, et al. (2023) Association analysis of mitochondrial genome polymorphisms with backfat thickness in pigs. *Animal biotechnology*, 35(1), 2272172.
- Rane R, et al. (2023) Complex multiple introductions drive fall armyworm invasions into Asia and Australia. *Scientific reports*, 13(1), 660.