

Resource Summary Report

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Stacks

RRID:SCR_003184

Type: Tool

Proper Citation

Stacks (RRID:SCR_003184)

Resource Information

URL: <http://creskolab.uoregon.edu/stacks/>

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Description: A software pipeline for building loci from short-read sequences, such as those generated on the Illumina platform. It was developed to work with restriction enzyme-based data, such as RAD-seq, for the purpose of building genetic maps and conducting population genomics and phylogeography.

Abbreviations: Stacks

Resource Type: data analysis software, data processing software, software resource, software application

Defining Citation: [PMID:23701397](#), [PMID:22384329](#), [DOI:10.1111/mec.12354](#)

Keywords: population genomics, genetic map, phylogenetics, genetics, next-generation sequencing, rad-seq, genotype-by-sequencing, bio.tools

Funding:

Availability: GNU General Public License, v3

Resource Name: Stacks

Resource ID: SCR_003184

Alternate IDs: OMICS_01567, biotools:stacks

Alternate URLs: <https://bio.tools/stacks>, <https://sources.debian.org/src/stacks/>

Record Creation Time: 20220129T080217+0000

Record Last Update: 20250423T060117+0000

Ratings and Alerts

No rating or validation information has been found for Stacks.

No alerts have been found for Stacks.

Data and Source Information

Source: [SciCrunch Registry](#)

Usage and Citation Metrics

We found 538 mentions in open access literature.

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

Dufresnes C, et al. (2025) Speciation and historical invasions of the Asian black-spined toad (*Duttaphrynus melanostictus*). *Nature communications*, 16(1), 298.

Patova A, et al. (2025) Population genomics and connectivity of *Vazella pourtalesii* sponge grounds of the northwest Atlantic with conservation implications of deep sea vulnerable marine ecosystems. *Scientific reports*, 15(1), 1540.

Amado D, et al. (2025) The genetic architecture of resistance to flubendiamide insecticide in *Helicoverpa armigera* (Hübner). *PloS one*, 20(1), e0318154.

Kniesz K, et al. (2025) High genomic connectivity within *Anatoma* at hydrothermal vents along the Central and Southeast Indian Ridge. *Scientific reports*, 15(1), 1971.

Domínguez M, et al. (2025) Genomics Reveal Population Structure and Intergeneric Hybridization in an Endangered South American Bird: Implications for Management and Conservation. *Ecology and evolution*, 15(1), e70820.

Torrado H, et al. (2025) Evolutionary Genomics of Two Co-occurring Congeneric Fore Reef Coral Species on Guam (Mariana Islands). *Genome biology and evolution*, 17(1).

Paris JR, et al. (2025) The Genomic Signature and Transcriptional Response of Metal Tolerance in Brown Trout Inhabiting Metal-Polluted Rivers. *Molecular ecology*, 34(1), e17591.

Ho DV, et al. (2024) Post-meiotic mechanism of facultative parthenogenesis in gonochoristic whiptail lizard species. *eLife*, 13.

Reatini B, et al. (2024) Chromosome-scale Reference Genome and RAD-based Genetic Map of Yellow Starthistle (*Centaurea solstitialis*) Reveal Putative Structural Variation and QTL Associated With Invader Traits. *Genome biology and evolution*, 16(12).

Worthington BM, et al. (2024) Serological evidence of sarbecovirus exposure along Sunda pangolin trafficking pathways. *BMC biology*, 22(1), 274.

Peralta DM, et al. (2024) A rapid approach for sex assignment by RAD-seq using a reference genome. *PloS one*, 19(4), e0297987.

Çetin C, et al. (2024) Population genetic structure in a self-compatible hermaphroditic snail is driven by drift independently of its contemporary mating system. *Ecology and evolution*, 14(8), e70162.

Raggi L, et al. (2024) Investigating the genetic basis of salt-tolerance in common bean: a genome-wide association study at the early vegetative stage. *Scientific reports*, 14(1), 5315.

Strait JT, et al. (2024) Local environments, not invasive hybridization, influence cardiac performance of native trout under acute thermal stress. *Evolutionary applications*, 17(2), e13663.

Taichi N, et al. (2024) Habitat diversification associated with urban development has a little effect on genetic structure in the annual native plant *Commelina communis* in an East Asian megacity. *Ecology and evolution*, 14(2), e10975.

Búci M, et al. (2024) Airports for the genetic rescue of a former agricultural pest. *Scientific reports*, 14(1), 17540.

Ramirez-Ramirez AR, et al. (2024) Comparing the performances of SSR and SNP markers for population analysis in *Theobroma cacao* L., as alternative approach to validate a new ddRADseq protocol for cacao genotyping. *PloS one*, 19(5), e0304753.

Brunharo CACG, et al. (2024) Chromosome-scale genome assembly of *Poa trivialis* and population genomics reveal widespread gene flow in a cool-season grass seed production system. *Plant direct*, 8(3), e575.

Yildirim Y, et al. (2024) Phylogeography and phenotypic wing shape variation in a damselfly across populations in Europe. *BMC ecology and evolution*, 24(1), 19.

Liu Y, et al. (2024) Conservation genetics and potential geographic distribution modeling of *Corybas taliensis*, a small 'sky Island' orchid species in China. *BMC plant biology*, 24(1), 11.