

# Resource Summary Report

Generated by [FDI Lab - SciCrunch.org](https://fdi-lab.sci-crunch.org) on Apr 15, 2025

## Bs-Seeker2

RRID:SCR\_020948

Type: Tool

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### Proper Citation

Bs-Seeker2 (RRID:SCR\_020948)

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### Resource Information

**URL:** [https://guoweilong.github.io/BS\\_Seeker2/index.html](https://guoweilong.github.io/BS_Seeker2/index.html)

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**Description:** Software tool as versatile aligning pipeline for bisulfite sequencing data. Used for mapping bisulfite sequencing data and generating DNA methylomes. Improves mappability over existing aligners by using local alignment. Maps reads from RRBS library by building special indexes with improved efficiency and accuracy. Provides additional function for filtering out reads with incomplete bisulfite conversion, which is useful in minimizing overestimation of DNA methylation levels.

**Synonyms:** Bisulfite Sequencing Seeker2, BS Seeker2

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

**Defining Citation:** [PMID:24206606](https://pubmed.ncbi.nlm.nih.gov/24206606/)

**Keywords:** Versatile aligning pipeline, bisulfite sequencing data, mapping bisulfite sequencing data, generating DNA methylomes, DNA methylation level, reads mapping, Reduced Represented Bisulfite Sequencing library, bio.tools

**Funding:** Institute of Genomics and Proteomics at UCLA ;  
NBRPC 2012CB316503;  
China Scholarship Council

**Availability:** Free, Available for download, Freely available

**Resource Name:** Bs-Seeker2

**Resource ID:** SCR\_020948

**Alternate IDs:** biotools:bs-seeker2

**Alternate URLs:** [http://pellegrini.mcdb.ucla.edu/BS\\_Seeker2/](http://pellegrini.mcdb.ucla.edu/BS_Seeker2/), <https://bio.tools/bs-seeker2>

**Record Creation Time:** 20220129T080353+0000

**Record Last Update:** 20250412T060305+0000

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## Ratings and Alerts

No rating or validation information has been found for Bs-Seeker2.

No alerts have been found for Bs-Seeker2.

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## Data and Source Information

**Source:** [SciCrunch Registry](#)

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## Usage and Citation Metrics

We found 2 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [FDI Lab - SciCrunch.org](#).

Lizamore D, et al. (2021) Elevated transcription of transposable elements is accompanied by het-siRNA-driven de novo DNA methylation in grapevine embryogenic callus. BMC genomics, 22(1), 676.

Bourguet P, et al. (2020) DNA polymerase epsilon is required for heterochromatin maintenance in Arabidopsis. Genome biology, 21(1), 283.