

# Resource Summary Report

Generated by [Kravitz](#) on Sep 10, 2026

## SVseq

RRID:SCR\_004804

Type: Tool

### Proper Citation

SVseq (RRID:SCR\_004804)

### Resource Information

**URL:** <http://www.engr.uconn.edu/~jiz08001/svseq2.html>

**Proper Citation:** SVseq (RRID:SCR\_004804)

**Description:** Software for accurate and efficient calling of structural variations with low-coverage sequence data. Version 2 uses the BAM files of paired Illumina reads with soft-clip signature as input. It calls both deletions and insertions.

**Abbreviations:** SVseq

**Synonyms:** SVseq2, SVseq1

**Resource Type:** software resource

**Defining Citation:** [PMID:22537045](#)

**Keywords:** structural variant, deletion, insertion, breakpoint, bio.tools

**Resource Name:** SVseq

**Resource ID:** SCR\_004804

**Alternate IDs:** OMICS\_00327, biotools:svseq

**Alternate URLs:** <https://bio.tools/svseq>

**Record Creation Time:** 20220129T080226+0000

**Record Last Update:** 20260905T132524+0000

### Ratings and Alerts

No rating or validation information has been found for SVseq.

No alerts have been found for SVseq.

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## Data and Source Information

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

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## Usage and Citation Metrics

We found 3 mentions in open access literature.

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

Qin S, et al. (2025) SVseq discloses the genomic complexity of different prenatal, de novo, apparently balanced chromosome rearrangements detected by CMA and karyotype. Italian journal of pediatrics, 51(1), 155.

Pirooznia M, et al. (2015) Whole-genome CNV analysis: advances in computational approaches. Frontiers in genetics, 6, 138.

Thangam M, et al. (2015) CRCDA--Comprehensive resources for cancer NGS data analysis. Database : the journal of biological databases and curation, 2015.