

# RSeqAn: Headers and wrappers for the SeqAn library in R

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## Summary

**RSeqAn** provides R with access to SeqAn (Döring et al. 2008; Reinert et al. 2017) header files. SeqAn is an open source C++ library of efficient algorithms and data structures for the analysis of sequences with a focus on biological data. It has been used for many popular bioinformatics tools, including Bowtie2 (Langmead and Salzberg 2012) and Tophat (Trapnell, Pachter, and Salzberg 2009). Many packages in R are sped up with C++ code: as of November 2018, out of 13525 packages on CRAN, 1493 of those use Rcpp (Eddelbuettel and Balamuta 2018). On Bioconductor (Huber et al. 2015), a repository specifically for bioinformatics packages, there is a similar proportion of packages using Rcpp: 150 out of 1649. However, to date these packages have not utilized SeqAn.

An R package that does aim to use SeqAn runs into two issues. First, R for Windows is built with the mingw compiler, which the SeqAn development team does not offer support for. Second, SeqAn implements several data types in order to make the library more efficient. Any data types in SeqAn must thus be extended with Rcpp through the templated functions `Rcpp::as` and `Rcpp::wrap`.

In **RSeqAn** v1.0 we provide access to SeqAn header files and add support for mingw so that it can be used in R for Windows as well. The development version also includes some wrappers to convert between R objects and SeqAn data types with a goal of eventually providing all necessary wrappers for easy interface with SeqAn. **RSeqAn** thus provides an easy way to write R bioinformatics packages that are much more efficient without need for the user to install SeqAn themselves. **RSeqAn** can be installed from Bioconductor, and documentation for it can be found in the package as well as online.

## Benchmarked Example

As a proof of concept for its utility, we have benchmarked a function (searching for adapter contamination) from **qckitfastq**, a package that uses **RSeqAn** for quality control on bioinformatics data compared to **ShortRead** (Morgan et al.

2009), another package on Bioconductor serving the same function that is purely written in R (Table 1). As can be seen, computing adapter content through `qckitfastq` is much faster than `ShortRead`.

Table 1: Benchmark results of `qckitfastq`, which uses `RSeqAn`, and `ShortRead`, which does not against a test file from each package.

File	Package	replications	elapsed	user.self	sys.self
<b>E-MTAB-1147</b>	ShortRead	100	13.3	11.58	1.37
	<code>qckitfastq</code>	100	5.159	4.902	0.091
<b>test.fq.gz</b>	ShortRead	100	6.612	5.739	0.567
	<code>qckitfastq</code>	100	0.195	0.105	0.019

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