

# libqcpp: A C++14 sequence quality control library

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#### **Software**

- Review 🗗
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# Summary

Libqcpp implements a variety of algorithms for Next-generation Sequencing (NGS) data quality control. These algorithms include:

- Sliding-window quality score trimming, using an algorithm based on Sickle (Joshi and Fass 2011).
- A combined adaptor removal and read merging algorithm for paired end reads that uses global pairwise alignment of reads. This algorithm is similar to AdapterRemoval (Lindgreen 2012).
- Cycle-wise summarisation of base quality scores, similar to FastQC (Andrews 2012)

Libqcpp allows simple composition of quality control pipelines that combine these features into a single unit. Application code can then simply read from a stream of sequence reads that have passed quality control measures. Optionally, parsing and quality control can occur in one or more background threads for efficiency. Reports detailing actions performed and summaries of results may be obtained in YAML format. Libqcpp includes trimit, a command line interface to these features for those not building their own applications.

Libqcpp uses the SeqAn library for sequence parsing and alignment (Döring et al. 2008), libyaml-cpp for YAML report generation, and Catch for unit testing. Documentation on API and command line usage is included, and available at https://qcpp.readthedocs.io/.

### References

Andrews, S. 2012. "FastQC A Quality Control Tool for High Throughput Sequence Data." http://www.bioinformatics.babraham.ac.uk/projects/fastqc/.

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Joshi, N A, and J N Fass. 2011. Sickle: A Sliding-Window, Adaptive, Quality-Based Trimming Tool for FastQ Files (version 1.33). https://github.com/najoshi/sickle.

Lindgreen, Stinus. 2012. "Adapter Removal: Easy Cleaning of Next-Generation Sequencing Reads."  $BMC\ Research\ Notes\ 5\colon 337.\ doi:10.1186/1756-0500-5-337.$