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## SeQuiLa-cov: A fast and scalable library for depth of coverage calculations --Manuscript Draft--

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Abstract:	Background: Depth of coverage calculation is an important and computationally intensive preprocessing step in a variety of next generation sequencing pipelines, including the analyses of RNA-seq data, detection of copy number variants, or quality control procedures.  Results: Building upon big data technologies, we have developed SeQuiLa-cov, an extension to the recently released SeQuiLa platform, which provides efficient depth of coverage calculations, reaching more than 100x speedup over the state-of-the-art tools. Performance and scalability of our solution allows for exome and genome-wide calculations running locally or on a cluster while hiding the complexity of the distributed computing with Structured Query Language Application Programming Interface.  Conclusions: SeQuiLa-cov provides significant performance gain in depth of coverage calculations streamlining the widely used bioinformatic processing pipelines.				
Corresponding Author:	Tomasz Gambin Warsaw University of Technology Warszawa, POLAND				
Corresponding Author Secondary Information:					
Corresponding Author's Institution:	Warsaw University of Technology				
Corresponding Author's Secondary Institution:					
First Author:	Marek Wiewiórka				
First Author Secondary Information:					
Order of Authors:	Marek Wiewiórka				
	Agnieszka Szmurło				
	Wiktor Kuśmirek				
	Tomasz Gambin				
Order of Authors Secondary Information:					
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	Dear Editor,				
	Thank you for accepting our manuscript for publication in GigaScience. Please, find below a detailed point-by-point response to all of your comments.				
	Most sincerely, Tomasz Gambin, Ph. D. ZSI-Bio group (http://biodatageeks.org)				

Institute of Computer Science Warsaw University of Technology E-mail: tgambin@ii.pw.edu.pl 1) Our data curators usually take an archival snapshot of the source code and test data, which will be posted via our repository GigaDB. Our curators will contact you shortly to prepare this dataset. Once this is ready, you will get a DOI for this GigaDB dataset. Please include a citation to your GigaDB dataset (including the DOI link) to your reference list, and cite this in the data availability section in the manuscript, where appropriate. Please follow this example format for the reference: [xx] Author1 N, Author2 N, AuthorX N. Supporting data [and code] for "Title of your manuscript". GigaScience Database. 2019. http://dx.doi.orgxxxxxxxxxxxxx We are looking forward to contacting us by data curators. 2) I note that all authors are also listed as "corresponding author" on the title page. Gigascience allows a maximum of two corresponding authors (see https://academic.oup.com/gigascience/pages/authorship\_guidelines ) - please discuss who (one or two authors) should take on this role. We double checked this and there is only one corresponding author (Tomasz Gambin) indicated on the title page. Please, let us know if you need us to change the way how the corresponding author is marked. 3) To avoid confusion during the production process, please remove any highlighting / tracking of changes that was made for the purpose of peer review. We removed all highlighting as requested. 4) Thank you for registering your tool at Scicrunch. The RRID should be mentioned as the last bullet point in the "software availability" section (below "licence"), in the following format: RRID: SCR NNNNN (see https://academic.oup.com/gigascience/pages/technical\_note) (the sentence regarding the RRID in the following section is then redundant and can be deleted). We created a new bullet point below "licence" and deleted the sentence in the following section as requested. Additional Information: Question Response Are you submitting this manuscript to a No special series or article collection? Experimental design and statistics Yes Full details of the experimental design and statistical methods used should be given in the Methods section, as detailed in our Minimum Standards Reporting Checklist. Information essential to interpreting the data presented should be made available in the figure legends. Have you included all the information requested in your manuscript?

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Manuscript in Preparation Paper

PAPER

# SeQuiLa-cov: A fast and scalable library for depth of coverage calculations

Marek Wiewiórka<sup>1,\*</sup>, Agnieszka Szmurło<sup>1,\*</sup>, Wiktor Kuśmirek<sup>1</sup> and Tomasz Gambin<sup>1,†</sup>

<sup>1</sup>Institute of Computer Science, Warsaw University of Technology, ul. Nowowiejska 15/19, 00-665 Warsaw, Poland,

#### **Abstract**

Background Depth of coverage calculation is an important and computationally intensive preprocessing step in a variety of next generation sequencing pipelines, including the analyses of RNA-seq data, detection of copy number variants, or quality control procedures. Results Building upon big data technologies, we have developed SeQuiLa-cov, an extension to the recently released SeQuiLa platform, which provides efficient depth of coverage calculations, reaching more than 100x speedup over the state-of-the-art tools. Performance and scalability of our solution allows for exome and genome-wide calculations running locally or on a cluster while hiding the complexity of the distributed computing with Structured Query Language Application Programming Interface. Conclusions SeQuiLa-cov provides significant performance gain in depth of coverage calculations streamlining the widely used bioinformatic processing pipelines.

**Key words**: NGS data analysis; depth of coverage; big data; distributed computing; SQL; CNV-calling; RNA-seq; quality control for sequencing data;

#### **Findings**

#### Introduction

Given a set of sequencing reads and a genomic contig, depth of coverage for a given position is defined as a total number of reads overlapping the locus.

The coverage calculation is a frequently performed but time-consuming step in the analysis of Next Generation Sequencing (NGS) data. In particular, Copy-Number Variant detection pipelines require obtaining sufficient read depth of the analyzed samples [1, 2, 3]. In other applications, the coverage is computed to assess the quality of the sequencing data (e.g. to calculate the percentage of genome with at least 30X read depth) or to identify genomic regions overlapped by insufficient number of reads for reliable variant calling [4]. Finally, depth of coverage is one of the most computationally intensive parts of differential expression analysis using RNA-seq data at

single-base resolution [5, 6, 7].

A number of tools supporting this operation have been developed, with 22 of them specified in Omictools catalog [8]. Well known, state-of-the-art solutions include: samtools depth [9], bedtools genomecov [10], GATK DepthOfCoverage [11], sambamba [12], and mosdepth [13] (see comparison presented in Table 1).

Traditionally, these methods calculate the depth of coverage using a pileup-based approach (introduced in samtools [9] and used in GATK [11]), which is inefficient since it iterates through each nucleotide position at every read in a Binary Alignment Map (BAM) file. An optimized, event-based approach has been proposed in bedtools [10] and mosdepth [13]. These algorithms use only specific 'events', i.e. start and end of the alignment blocks within each read (Figure 1A) instead of analyzing every base of each read, which significantly reduces the overall computational complexity.

Samtools and bedtools depth of coverage modules do not

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<sup>\*</sup>Contributed equally.

<sup>†</sup>Corresponding author.

#### **Key Points**

- $\cdot$  SeQuiLa-cov allows for high-coverage ( $\sim$ 60x) genome-wide depth of coverage calculations in less than one minute.
- SeQuiLa-cov provides ANSI SQL compliant API for accessing and analyzing of aligned sequencing reads data.

Table 1. Comparison of leading coverage calculation software tools.

Functionality			Implementation					
tool	approach	bases	blocks	windows	language	Intel GKL	parallelism type	interface
samtools	pileup	yes	no	no	С	no	none	cmd line
bedtools	events	yes	yes	no	C++	no	none	cmd line
GATK <sup>1</sup>	pileup	yes	no	no	Java	yes	distributed	cmd line
sambamba	pileup	no	yes	yes	D	no	multithreaded	cmd line
mosdepth	events	no	yes	yes	Nim	no	multithreaded <sup>2</sup>	cmd line
SeQuiLa-cov	events	yes	yes	yes	Scala	yes	distributed	Scala, SQL

<sup>&</sup>lt;sup>1</sup>GATK DepthOfCoverage has not yet been ported to the latest version, i.e. GATK 4.x

provide any support for multi-core environment. Mosdepth implements parallel BAM decompression, but its main algorithm remains sequential. Sambamba, on the other hand, promotes itself as a highly parallel tool, implementing depth of coverage calculations in a map-reduce fashion utilizing multiple threads on a single node. Regardless of parallelization degree, all of the above mentioned tools share a common bottleneck caused by using a single thread for returning results. Finally, GATK was the first genomic framework providing a support for distributed computations, however, the DepthOfCoverage method has not been ported yet to the current software release of the toolkit.

We present the first fully scalable, distributed, SQLoriented solution designated for the depth of coverage calculations. SeQuiLa-cov, an extension to the recently released Se-QuiLa [14] platform, runs a redesigned event-based algorithm for the distributed environment and provides convenient, SQLcompliant interface.

#### Algorithm and implementation

Consider input data set, read\_set, of aligned sequencing reads sorted by genomic position from a BAM file partitioned into the n data slices (read\_set<sub>1</sub>, read\_set<sub>2</sub>, ..., read\_set<sub>n</sub>) (Figure 1B).

In the most general case, the algorithm can be used in a distributed environment where each cluster node computes the coverage for the subset of data slices using the event-based method. Specifically, for the i-th partition containing the set of reads ( $read\_set_i$ ), the set of  $events_{i,chr}$  vectors (where chr is an index of genomic contig represented in read\_set) is allocated and updated, based on the items from read\_set<sub>i</sub>. For all reads, the algorithm parses the CIGAR string and for each continuous alignment block characterized by start position and length len it increments by one the  $events_{i,chr}(start)$  and decrements by one the value of  $events_{i,chr}(start + len)$ . To compute the partial coverage vector for partition i and contig chr, a vector value at the index *j* is calculated as follows:

$$partial\_coverage_{i,chr}(j) = \sum_{m=1}^{J} events_{i,chr}(m).$$

The result of this stage is a set of partial\_coverage<sub>i,chr</sub> vectors distributed among the computation nodes. To calculate the final coverage for the whole read\_set, an additional step of correction for overlaps between the partitions is required. An overlap  $overlap_{i,chr}$  of length l between vectors partial\_coverage<sub>i,chr</sub> and partial\_coverage<sub>i+1,chr</sub> may occur on the partition boundaries where l tailing genomic positions of

partial\_coverage<sub>i,chr</sub> are the same as l heading genomic positions of  $partial\_coverage_{i+1,chr}$  (see Figure 1C).

If an overlap is identified then the coverage values from the  $partial\_coverage_{i,chr}$ 's l-length tail are added into the  $partial\_coverage_{i+1,chr}$ 's head and subsequently the last l elements of partial\_coveragei,chr are removed. Once this correction step is completed, non-overlapping  $coverage_{i,chr}$  vectors are collected and yield the final coverage values for the whole

The main characteristic of the described algorithm is its ability to distribute data and calculations (such as BAM decompression and main coverage procedure) among the available computation nodes. Moreover, instead of simply performing full data reduction stage of the partial coverage vectors, our solution minimizes required data shuffling among cluster nodes by limiting it to the overlapping part of coverage vectors. Importantly, SeQuiLa-cov computation model supports fine-grained parallelism at user-defined partition size in contrary to the traditional, coarse-grained parallelization strategies that involve splitting input data at a contig level.

#### **Implementation**

We have implemented SeQuiLa-cov in Scala programming language using the Apache Spark framework. To efficiently access the data from a BAM file we have prepared a custom data source using Data Source Application Programming Interface (API) exposed by SparkSQL. Performance of the read operation benefits from the Intel Genomics Kernel Library (GKL) [15] used for decompressing the BAM files chunks and from predicate pushdown mechanism that filters out data at the earliest stage.

The implementation of the core coverage calculation algorithm aimed at minimizing, whenever possible memory footprint by using parsimonious data types, e.g. Short type instead of Integer, and efficient memory allocation strategy for large data structures, e.g. favoring static Arrays over dynamic size ArrayBuffers. Additionally, to reduce the overhead of data shuffling between the worker nodes in the correction for the overlaps stage we used Spark's shared variables [16] accumulators and broadcast variables (Figure 1C). Accumulator is used to gather information about the worker nodes' coverage vector ranges and coverage vector tail values, that are subsequently read and processed by the driver. This information is then used to construct a broadcast variable distributed to the worker nodes in order to perform adequate trimming and summing operations on partial coverage vectors.

<sup>&</sup>lt;sup>2</sup>Only for BAM decompression

data operation type cores samtools bedtools sambamba mosdepth SeQuiLa-cov 1 2h 14m 58s 1 10h 41m 27s 2h 44m 0s 1h 46m 27s 1h 47m 5s blocks 5 2h 47m 53s 36m 13s 26m 59s 10 2h 50m 47s 34m 34s 13m 54s WGS 1 1h 46m 50s 1h 22m 49s 1h 24m 8s fixed-length windows 1h 41m 23s 20m 3s 18m 43s 5 10 1h 50m 35s 17m 49s 9m 14s 12m 26s 1 23m 25s 25m 42s 6m 43s 6m 54s blocks 5 25m 46s 2m 25s 1m 47s 10 25m 49s 2m 20s 1m 4s WES 1 14m 36s 6m 11s 6m 29s fixed-length windows 5 14m 54s 2m 8s 1m 42s 10 14m 40s 2m 14s 1m 1s

Table 2. Benchmarking leading solutions against SeQuiLa-cov on WES/WGS data in performing blocks and windows calculations

Both samtools and bedtools calculate coverage using only a single thread, however, their results differ significantly, with samtools being around twice as fast. Sambamba positions itself as a multithreaded solution although our tests revealed that its execution time is nearly constant, regardless of the number of CPU cores used, and even twice as slow as samtools. Mosdepth achieved speedup against samtools in blocks coverage and against sambamba in windows coverage calculations, however, its scalability reaches limit at 5 CPU cores. Finally, SeQuiLa-cov, achieves nearly identical performance as mosdepth for the single core but the execution time decreases substantially for greater number of available computing resources which makes this solution the fastest when run on multiple cores and nodes.

<sup>1</sup>per-base results are treated as blocks output. Samtools lacks the functionality of blocks coverage calculations, however, we included this tool in our benchmark for completeness, treating its per-base results as blocks outcome assuming that both result types require nearly the same resources.

#### **Functionality**

#### Supported coverage result types

SeQuiLa-cov features three distinct result types: per-base, blocks, and fixed-length windows coverage (Figure 1A). For perbase, the depth of coverage is calculated and returned for each genomic position making it the most verbose output option. The method producing block level coverage (blocks) involves merging adjacent genomic positions with equal coverage values into genomic intervals. As a consequence, fewer records than in case of per-base output type are generated with no information loss. The *fixed-length windows* the algorithm generates set of fixed length, tiling, non-overlapping genomic intervals and returns arithmetic mean of coverage values over positions within each window.

#### ANSI SQL compliance

SeQuiLa-cov solution promotes SQL as a data query and manipulation language in genomic analysis. Data flows are performed in SQL-like manner through the custom data source supporting convenient Create Table as Select and Insert as Select methods. SeQuiLa-cov provides a table abstraction over existing alignment files, with no need of data conversion, which can be further queried and manipulated in a declarative way. The coverage calculation function *bdg\_coverage*, as described in Algorithm sub-section, has been implemented as table-valued function(Figure 1D).

#### **Execution and integration options**

SeQuiLa-cov can be used as an extension to Apache Spark in a form of external JAR dependency or can be executed from command-line as a Docker container. Both options can be run locally (on a single node) or on a Hadoop cluster using Yet Another Resource Negotiator (YARN). (See Project Documentation for sample commands). The tool accepts BAM/CRAM files as input and supports processing of short and long reads. The tabular output of the coverage computations can be stored in various file formats, e.g. binary (ORC, Parquet), as well as text (CSV, TSV). The tool can be integrated with state-of-the-art applications through text files, or can be used directly as an additional library in bioinformatics pipelines implemented in Scala, R, or Python.

#### **Benchmarking**

We have benchmarked SeQuiLa-cov solution with leading software for depth of coverage calculations, specifically samtools depth, bedtools genomeCov, sambamba depth and mosdepth (results of DepthOfCOverage from outdated GATK version are available in supplementary data). The tests were performed on the aligned WES and WGS reads from the NA12878 sample (see Methods for details) and aimed at calculating blocks and window coverage. To compare the performance and scalability of each solution, we have executed calculations for 1, 5, and 10 cores on a single computation node (see Table 2).

Samtools depth and bedtools genomeCov are both natively non-scalable and were run on a single thread only. Exomewide calculations exceeded 10 minutes and genome-wide analyses took over two hours in case of samtools, while bedtools' performance was significantly worse, i.e ~1.9x for WES and ~4.75x for WGS. Sambamba depth declares to take advantage of fully parallelized data processing with the use of multithreading. However, our results revealed that even when additional threads were used, the total execution time of coverage calculations remained nearly constant and greater than samtools's result. Mosdepth shows significant speedup (~1.3x) against samtools when using single thread. This performance gain increases to ~3.7x when using 5 decompression threads, however, it does not benefit from adding additional CPU power. In case of fixed-length window coverage mosdepth achieves over ~1.3 speedup against sambamba.

SeQuiLa-cov achieves performance similar to mosdepth when run using a single core. However, SeQuiLa-cov is ~1.3x and ~2.5x as fast as mosdepth when using 5 and 10 CPU cores, respectively, demonstrates its better scalability. The similar performance characteristic is observed for both blocks and fixed-length windows methods.

To fully assess the scalability profile of our solution, we have performed additional tests in a cluster environment (see Methods for details). Our results show that when utilizing additional resources (i.e. more than 10 CPU cores), SeQuiLa-cov is able to reduce the total computation time to 15 seconds for WES and less than one minute for WGS data (Figure 2). Scalability limit is achieved for 200 and ~500 CPU cores in case of WES and WGS data, respectively.

To evaluate the impact of Intel GKL library on deflate operation (BAM bzgf block decompression), we have performed blocks coverage calculations on WES data on 50 CPU cores. The

results showed on average ~1.18x speedup when running with Intel GKL deflate implementation.

Figure 1. SeQuiLa-cov: functionality, algorithm and implementation

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partial coverage 1

Panel A shows the general concept of events-based algorithm for depth of coverage calculation. Given a genomic chromosome and a set of aligned sequencing reads, the algorithm allocates events vector. Subsequently, it iterates the list of reads and increments/decrements by one the values of the events vector at the indexes corresponding to start/end positions of each read. The depth of coverage for a genomic locus is calculated using the cumulative sum of all elements in the events vector preceding specified position. The algorithm may produce three typically used coverage types: (i) per-base coverage, which includes the coverage value for each genomic position separately, (ii) blocks which lists adjacent positions with equal coverage values are merged into single interval, and (iii) fixed-length windows coverage that generates set of equal-size, non-overlapping and tiling genomic ranges and outputs arithmetic mean of base coverage values for each region.

ПППП

ПППП

final\_coverage\_n

4↓

partial coverage

Panel B presents the provided SQL API to interact with NGS data. The first statement creates a relational table read set over compressed BAM files using the provided custom Data Source, whereas the second statement demonstrates the use of bdg\_coverage function to calculate depth of coverage for a specified sample. The presented call for coverage method takes sample identifier (sample1) and result type (blocks) as input parameters. bdg\_coverage is implemented as a tablevalued function. Therefore, it outputs a table as a result allowing for customizing a query using Data Manipulation Language e.g. in the SELECT or WHERE clause. For the purpose of this example, we assume that BAM file for sample1 contains only reads from chr3.

Panel C shows the concept of distributed version of events-based algorithm. Assuming that we run our calculations in a distributed environment, the computation  $nodes \ do \ not \ work \ on \ the \ whole \ input \ data \ set \ (table \ \textit{read\_set}) \ but \ on \ \textit{n} \ smaller \ data \ partitions \ (\textit{slice}_1, \ \textit{slice}_2, \ \dots \ \textit{,slice}_n), \ each \ containing \ subset \ of \ input \ aligned$ reads. First the algorithm calculates partial events vector for available data slices and subsequently produces corresponding partial partial\_coverage vector. Due to the possibility of overlapping of ranges between two consecutive data slices, additional correction step needs to be performed. When an overlap is identified, the corresponding coverage values from the preceding vector's tail are cut and added to the head values of the subsequent vector. On the figure two overlaps were shown, one of them situated between partial\_coverage1 and partial\_coverage2 (overlap12 of length 4) encompassing positions chr3:101-104. The coverage values from partial\_coverage1 for overlap12 are removed from partial\_coverage1 and added to the head of partial\_coverage2. As a result, a set of non-overlapping coverage vectors are calculated, which is further integrated into the depth of coverage for the whole input data set.

Panel D presents the implementation details of SeQuiLa-cov. We have used the Apache Spark environment, where a single driver node runs the high-level driver program, which schedules tasks for multiple worker nodes. On each worker node, a set of data partitions are accessed and manipulated in order to generate events and partial\_coverage vectors. To gather data about partial\_coverage vectors' ranges along with tailing coverage values, and to distribute data needed for rearranging coverage vector values and ranges, we have used Spark's shared variables accumulator and broadcast, respectively.

Finally, our comprehensive functional unit testing showed that results calculated by SeQuiLa-cov and samtools depth are identical.

#### **Conclusions**

The application of the recent advancements in big data technologies and distributed computing can contribute to both speeding up genomic data processing and management. Analysis of large genomic data sets require efficient, accurate, and scalable algorithms to perform calculations utilizing the computing power of multiple cluster nodes. In this work, we show that with sufficiently large cluster genome-wide coverage calculations may last less than a minute and at the same time being over 100x faster than the best single-threaded solution.

Although the tool can be integrated with non-distributed software, our primary aim is to support large scale processing pipelines and the full advantage of SeQuiLa-cov's scalability and performance will be available once it is deployed and executed in a distributed environment. We expect that there will be a growing number of scalable solutions (Big Data Genomics project [17] with tools DECA and Cannoli as well as GATK4 (https://software.broadinstitute.org/gatk/gatk4)) that can take advantage of reading input data directly from distributed storage systems.

SeQuiLa-cov is one of the building blocks of SeQuiLa [14] ecosystem, which initiated the move towards efficient, distributed processing of genomic data and providing SQLoriented API for convenient and elastic querying. We foresee that following this direction will enable the evolution of genomic data analysis from the file-oriented to table-oriented processing.

#### **Methods**

#### Test data

We have tested our solution using reads from NA12878 sample which were aligned to hg18 genome. WES data containing over 161 million of reads weights 17 GB and WGS data include over 2,6 billion of reads taking 272 GB of disk space. Both BAM files were compressed at the default BAM's compression level (5).

#### **Testing environment**

To perform comprehensive performance evaluation, we have setup a test cluster consisting of 28 Hadoop nodes (1 edge node, 3 master nodes and 24 data nodes) with Hortonworks Data Platform 3.0.1 installed. Each data node has 28 cores (56 with hyper-threading) and 512 GB of RAM, YARN resource pool has been configured with 2640 virtual cores and 9671 GB RAM.

#### **Investigated solutions**

In our benchmark we have used the most recent versions of the investigated tools i.e. samtools version 1.9, bedtools 2.27.0, sambamba 0.6.8, mosdepth version 0.2.3 and SeQuiLa-cov version 0.5.1.

#### Availability of source code and requirements

- · Project name: SeQuiLa-cov
- Project home page: http://biodatageeks.org/sequila/
- Source code repository: https://github.com/ZSI-Bio/

#### bdg-sequila

· Operating system: Platform independent

· Programming language: Scala · Other requirements: Docker · License: Apache License 2.0

• RRID: SCR\_017220

#### Availability of supporting data and materials

The Docker image is available at https://hub.docker.com/r/ biodatageeks/. Supplementary information on benchmarking procedure as well as test data are publicly accessible at project documentation site http://biodatageeks.org/sequila/ benchmarking/benchmarking.html#depth-of-coverage.

#### **Declarations**

#### List of abbreviations

API - Application Programming Interface

BAM - Binary Alignment Map

GKL - Genomics Kernel Library

NGS - Next Generation Sequencing

SQL - Structured Query Language

YARN - Yet Another Resource Negotiator

WES - Whole Exome Sequencing

WGS - Whole Genome Sequencing

#### Consent for publication

Not applicable

#### **Competing interests**

None of the authors have any competing interests.

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#### **Author's Contributions**

MW - conceptualization, formal analysis, investigation, software and writing. AS - data curation, formal analysis, investigation, software, visualization and writing. WK - formal analysis, investigation, writing. TG - formal analysis, supervision, investigation, visualization and writing. All authors approved the final manuscript.

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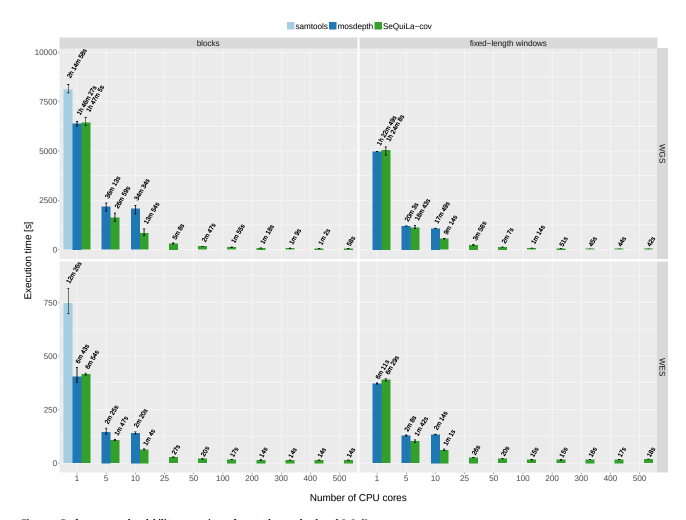


Figure 2. Performance and scalability comparison of samtools, mosdepth and SeQuiLa-cov Each experiment setting was repeated several times and the height of each bar along with the corresponding error bars indicate the average, as well as the minimum  $and \ maximum \ execution \ time, \ respectively. \ The \ best \ pileup-based \ solution \ is \ definitely \ slower \ (two \ times \ for \ WGS \ calculations) \ than \ both \ event-based \ solutions$ what clearly shows the superiority of the latter one. Mosdepth execution time scales up to 5 cores, afterwards it shows no furthe gain in performance. SeQuiLa-cov has nearly the same execution time results as mosdepth for both blocks and windows calculations for a single core, but scales out desirably utilizing all 500 CPU cores on cluster nodes and at the same time performing WGS calculations in less than 1 minute.

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