A quantitative assessment of the Hadoop framework for analysing massively parallel DNA sequencing data (Hadoop Or Not Hadoop)

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Abstract

New high-throughput technologies such as massively parallel sequencing has transformed the life sciences into a data-intensive field. With increasing data volumes comes the necessity to analyse data in parallel using high-performance computing resources, but doing this effectively can be laborious and challenging. Hadoop, emerging in the last decade, is a framework that automatically distributes data and computation and has been shown to scale to thousands of nodes. Herein we report a quantitative comparison of Hadoop to regular high-performance computing resources for aligning short reads and calling variants for five datasets of different sizes up to 250 gigabases. In order to increase performance of existing software and obtain a better comparison we modified and wrote new analysis scripts. From the observered scaling relations we are able to draw conclusions about the perspectives of the approaches, leading to the conclusion that as data set sizes reach 100 gigabases, the Hadoop-based pipelines become performance-competitive with a canonical high-performance cluster solution. As data sets in biological sequencing are sure to increase with time, Hadoop and similar frameworks are very interesting technologies that we envision will play a key role in the future of biological data analysis.

1 Introduction

Since its inception, massively parallel DNA sequencing, also referred to as Next Generation Sequencing (NGS) technology, has been an extremely bountiful source of data giving insight into the workings of biological machinery [?, ?]. Decreasing sequencing costs facilitates and promotes larger and larger studies with increasingly larger data sizes, and extracting useful information from these voluminous amounts of data is transforming biology into a data-intensive discipline. As an example of the scale of the demands, consider that a single Illumina high-throughput sequencing run produces approximately 3 TB of raw data in 10 days [?].

A common step of NGS data analysis consists of mapping short reads to a reference sequence and then finding the genetic variations specific to the sample. Many

of the most widespread tools for this purpuse, like BWA [1], Bowtie [2] and Samtools [3], are not made with distributed computing in mind. Many others do not even have the native ability to use multiple cores.

The most common approach to speed up NGS tools is to parallelise within a compute node (multi-core parallelisation) using shared memory parallelism (OMP) [4], but this approach is naturally limited by the number of cores per node, which does not usually exceed 16. confirm this! For the tools which do not support OMP natively, e.g. Samtools, variant calling can be parallelised by creating a separate process for each chromosome, or using GNU Parallel Linux utility [5]. Of great importance is that a multi-core approach does not improve the performance of operations that are limited by local disk or network throughputs, whereas splitting the dataset and multi-node parallelisation generally are not bound by these constraints.

Message Passing Interface (MPI) [6] is a common way to implement multi-node parallelisation, but writing efficient MPI-programs for hundreds of cores is a non-trivial task since thread synchronisation (or load balancing) has to be woven into the software code and there are few existing solutions available for processing sequencing data [7, 8, 9].

Another common way to introduce parallelisation to NGS analysis pipelines in Linux systems is to use Bash scripting. This involves using existing utilities and cluster tools to split the data into chunks, process them on the separate nodes, and merge the results afterwards. This kind of solution benefits from both MPI-like and OMP parallelisation and provides good performance, but the development requires substantial expertise in order to be efficient. Since the process is tightly coupled to the local computational cluster and network architectures, it might also not be possible to be re-used in other settings.

The Map-Reduce (MR) programming paradigm [10] offers a compelling alternative for running tasks in a massively parallel way. This paradigm, however, shifts the focus from the best performance to scalability, suited for managing huge datasets of sizes up to several terabytes [11]. The most prevalent open source implementation of Map-Reduce is Hadoop [10, 12]. Fix bib error (can't have both editor and author for this ref type) The Hadoop MR framework provides automatic distribution of computations over many nodes as well as automatic failure recovery (failure of individual jobs or computing nodes by storing multiple copies on different nodes), and automated collection of results [12]. Hadoop Distributed File System (HDFS) is a complementary component that stores data by automatically distributing it over the entire cluster, writing data blocks onto the local disk of each node and therefore effectively moving the computation to the data and reducing network traffic. HDFS provides a storage system whose bandwidth and size scales with the number of nodes in the cluster [13], which is very different from the properties of the usual HPC cluster network architecture.

In this manuscript we focus on the question when that's two questions Hadoop is an appealing alternative to the program tools generally found in HPC centers for DNA-seq analysis. Since Hadoop is written in Java, which is slower than the standard HPC programming languages like C or Fortran, we seek to estimate an average data size when it starts to be worthwhile to use Hadoop from a performance perspective. We use five datasets with short reads of different sizes, align them against the appropriate reference genome, and call the variances. For the existing Hadoop software we propose modifications, in order to benefit fully from massively parallel nature of MR computations. For the classical HPC DNA-seq analysis programs we

developed a set of Bash scripts utilizing multiple nodes for short read alignment and exploiting the network fully, thereby speeding up calculations. The execution times for each dataset were collected and scaling relations were analysed to answer the question in focus.

2 Methods

2.1 Datasets

We used publicly available DNA-seq datasets (I–III) as well as a synthetic dataset (IV) of *A.thaliana*, the well-known model plant, and a dataset (V) of two *H.sapiens* individuals (Table 1). Data for datasets I-III and V were obtained using Illumina/HiSeq sequencing platforms. Further information about the datasets is provided in the Supplementary material section.

Table 1: Datasets used in the comparison.

dataset	organism	size in Gbases
I	A.thaliana	1.4
II	A.thaliana	7.0
III	A.thaliana	30.0
IV	A.thaliana, the artificial dataset created using Samtools package	100.0
V	H.sapiens, two individuals (GM12750 and GM12004), sample SRR499924	250.0

2.2 Analysis pipelines

We constructed two pipelines for identifying single-nucleotide polymorphisms (SNPs) from short read data, one based on Hadoop and the other on regular HPC with a batch processing system (hereafter referred to as HPC). Our experiments then consisted of running the pipelines for the selected input dataset and measuring the wall-clock run time for each pipeline stage. All experiments were repeated several times and the results averaged considering the variations in table 4, why not use medians to obtain a data point for the particular combination of data size and computational platform. Acknowledging that there are different approaches and software for conducting bioinformatic analysis for HPC (e.g., GATK [14]), we decided to create the analysis pipeline as simple as possible to be able to make a better comparison with the one implemented for Hadoop.

1. HPC approach

Short read alignment: Bowtie ver. 0.12.8 SNP calling: Samtools ver. 0.1.19

2. Hadoop approach: Crossbow [15]

Short read alignment: Bowtie ver. 0.12.8

SNP calling: SOAPsnp 1.02 [16]

In the HPC pipeline, reads were aligned with Bowtie, followed by sorting the mapped reads and SNP calling with Samtools. The Bowtie aligner natively implements OMP, meaning that with 8 cores on the same computer the result can be

theoretically obtained 8 times faster than on a single core. Likewise, Samtools (as of version 0.1.19) also offers shared memory parallelism for several of its functions. Where available these features were used to improve the analysis speed. The exact workflow used is available in github? the code repository created for this work [?]. missing ref

The equivalent Hadoop-based pipeline was implemented with Crossbow. The input data and the indexed genome reference were copied to Hadoop's storage system (HDFS) before starting the experiments. Crossbow implements a short pipeline that pre-processes the input data, transforming it into a format suitable for the alignment stage, and then continues to use Bowtie for alignment and SoapSNP to call SNPs. Unfortunately, Crossbow's preprocessor is not written in MR manner, and thus cannot be run in a massively parallel way. Due to this limitation, this basic step threatened to be the most time-consuming procedure in our test pipeline and bias our experiments. To overcome this bottleneck we substituted Crossbow's preprocessor with our own MR implementation (available in github? the code repository [?]).

2.3 Computational resources

To run the HPC analysis pipeline we used a computational cluster at UPPMAX, which is heavily used for analysis of NGS-data [17], on nodes equipped with 8 dual-core CPUs. Data and reference genomes were stored on a parallel, shared-storage system. The Hadoop test platform was deployed on a private cloud at UPPMAX using the OpenNebula [18] cloud manager. The cluster was set up with Cloudera Hadoop distribution version 2.0.0-mr1-cdh4.5.0 [19]. For details on computational resources, see Supplementary material.

3 Results

3.1 Modified preprocessing stage

A native preprocessing stage in Crossbow provides great flexibility in delivering the data to the Hadoop cluster, where data can be downloaded from Amazon S3, FTP and HTTP servers over the Internet [15]. The only way to introduce parallelisations in the native preprocessing stage is to split the read files into smaller chunks and to generate a manifest file listing all these chunks. This, however, is lacking the massively parallel way of treating the data.

Assuming that sequencing platforms are usually in close proximity with facilities that provide both storage and computers, we considered that the data already had been downloaded to storage that Hadoop could access. We rewrote the Crossbow preprocessing stage scripts in a MR-manner, benefiting from its massively parallel nature. The requirement is that the FASTQ data are BZIP2 archived and accessible by Hadoop. For our case the storage with the data was mounted with SSHFS to one of the Hadoop nodes, from where the BZIP2'ed data were transferred to the HDFS storage. BZIP2 provides very good compression and is splittable, meaning that the archive can be expanded in a parallel manner. Also, BZIP2 format is natively supported by Hadoop, i.e., there is no difference in dealing with the FASTQ data or with its BZIP2 archive.

Table 2: Timings (in minutes) with Crossbow's native read preprocessor and with derived approach for datasets II, III, IV on a 56 cores Hadoop cluster, using the Crossbow, and with the BASH script on a single HPC node with 16 cores.

Dataset	II (7 Gbases)	III (30 Gbases)	IV (100 Gbases)
Crossbow native	60.6 ± 0.7	299.0 ± 2.3	673 ± 1.0
Hadoop, this work	7.0 ± 0.0	20.7 ± 0.4	52.4 ± 0.1
BASH, this work	7.4 ± 0.0	31.1 ± 0.1	114.5 ± 0.3

The Hadoop streaming library offers a possibility to write Mapper and Reducer Introduce in introduction for Hadoop jobs in any programming language. Our Python scripts efficiently process short reads, produced by Illumina sequencing platform of different versions, as well as the FASTQ files converted from SRA format (NCBI Sequence Read Archive).ref? The problem of lacking the unique FASTQ header standard was solved by the rewriting the header based on a SHA-1 hash function, and the reads mates were labeled with ".1" and ".2" suffixes. To ensure that both the forward and reverse reads of the same pair would end up on the same Reducer, secondary Mapper key-sorting mechanisms were involved.

In order to compare the native preprocessor and the proposed one, the data was put on HDFS beforehand to eliminate possible network delays, e.g., while downloading data from Amazon. Table 2 illustrates the benefits of our approach. To complete the comparison the same preprocessing was performed with a BASH script for the same data located on the HPC storage. The script (see Supplementary information) ran on a single HPC node, utilizes multiple cores, with the limiting factor being the hard disk IO performance.

3.2 Accuracy of pipelines

Since our HPC and Hadoop approaches use different SNP callers (Samtools and SOAPsnp, correspondingly) we should not expect them to deliver perfectly matching SNP lists, but still we expect them to capture and correctly identify the mutations. We tested the correctness just for the smallest dataset (I). The mutation $C \rightarrow T$ on chromosome 4 at position 16702262[20] was successfully localized by both applications.

3.3 Scalability of HPC and Hadoop approaches

To demonstrate the scalability of the HPC approach we collected running times for dataset I as a function of the number of cores used (Table 3). Consider replacing with a graph. The aligning mapping written in table, be consistent process in Bowtie scales fairly well, but the SNP calling part in Samtools is a bottleneck, scaling worse than Bowtie, and consuming a progressively larger portion of the total calculation time even after parallelising analysis by chromosome. For the given datasets (I–V) the timings for alignment against the corresponding genomes and SNP calling for the HPC and Hadoop approaches were collected (Table 4).

One of the attractive sides of using Hadoop MR is its almost linear scalability, i.e., calculation time linearly depends on the size of the dataset[15, 21], regardless

Table 3: Calculation time for Dataset I executed on different number of cores on a node of HPC cluster. awkward title, consider replacing with graph (and new title)

N cores	timi	speed-ups				
	mapping	SNP calling	total	total	map	SNP call
1	22.75 ± 1.24	$26.1 {\pm} 0.7$	48.9	1.00	1.00	1.00
2	11.00 ± 0.05	16.9 ± 1.3	27.9	0.88	1.03	0.77
4	5.91 ± 0.04	11.7 ± 0.5	17.6	0.69	0.96	0.56
6	$4.24{\pm}0.04$	$8.6 {\pm} 0.8$	12.8	0.64	0.89	0.51
8	3.44 ± 0.03	8.6 ± 0.9	12.1	0.51	0.83	0.38
10	$2.91 {\pm} 0.01$	7.7 ± 0.5	10.6	0.46	0.78	0.34
12	2.64 ± 0.04	7.5 ± 0.5	10.1	0.40	0.72	0.29
14	$2.57{\pm}0.03$	7.5 ± 0.6	10.1	0.35	0.63	0.25
16	2.88 ± 0.06	7.4 ± 0.4	10.2	0.30	0.49	0.22

Table 4: Timings (in minutes) for HPC and Hadoop deployments for different dataset sizes. Dataset is shown in brackets by roman numerals, "f.r." stands for "forward reads". Big deviation what aspect of deviation, not fixable? for the HPC is due to Samtools BAM sorting, which is IO and memory intensive, thus strongly depend on the queuing at the HPC cluster. No p= used here.

data, Gbases	1.4	3.5	7.0	15.0	30.0	100.0	250
	(I)	(II, f.r.)	(II)	(III, f.r.)	(III)	(IV)	(V)
Hadoop, 56 cores	_	_	39	62	108	250	1125
Hadoop, 56 cores	18 ± 0	18 ± 0	29 ± 0	47 ± 0	89 ± 0	238 ± 1	1164 ± 14
HPC1, 8 cores	41	96	157	307	596	1490	_
HPC, 16 cores	17 ± 1	22 ± 6	43 ± 3	81 ± 9	172 ± 15	467 ± 60	> 48 hours

the scaling nature of the underlying program (Samtools, Bowtie, etc.). Figure 1, based on Table 4, shows the calculation time as a function of the dataset size for the 56 cores Hadoop cluster and datasets I-IV.

3.4 Comparing Hadoop and HPC runtime efficiency

Hadoop was designed to digest huge datasets[10, 11]. One can propose then that the larger the dataset is, the more suitable Hadoop becomes compared to the HPC approach. In order to compare the "suitability" for different types of calculation platforms (Hadoop and HPC) each being run on a different number of cores, we constructed the following function:

$$F = T_p \times p$$
,

where T_p is the calculation time on p cores. This trivial/obvious formula doesn't deserve such special treatment.

Using the data from Table 4, we plot the ratio F_{Hadoop}/F_{HPC} as a function of the reciprocal dataset size (Figure 2). Extrapolation to zero on the X-axis estimates

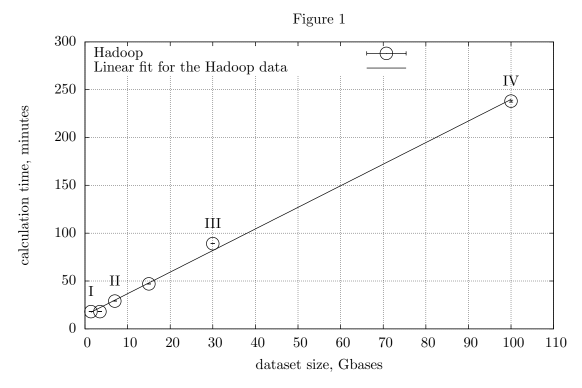


Figure 1: Calculation time for selected datasets (labeled with roman numerals) for the 56 cores Hadoop cluster shows a linear scaling with dataset size.

the ratio for the hypothetical infinite dataset. As one can see, the Hadoop approach becomes more and more effective compared to the HPC scenario as the dataset size increases. The parabolic extrapolation for the ratio is 1.70 ± 0.01 , meaning that Hadoop running even in the virtualized environment of a private cloud assembled on moderate hardware, is competitive with the HPC approach run on "bare metal" of modern hardware for datasets greater than 100 Gbases (dataset IV), which is typical for human genome sequencing with a sequencing depth of about 30x.

The simplest curve to fit the points nicely is a parabolic function, importantly not a linear one. Considering the linear scaling for the Hadoop approach for the same datasets, we conclude that HPC approach scales worse than linearly. We suspect the main reason for such a behavior is the fact that for large datasets (dataset III-V in our case) the SNP calling routine with Samtools becomes more memory and time demanding than the actual mapping with Bowtie. If lacking RAM, Samtools implementation swaps to disk, creating multiple (up to hundreds) temporary files while sorting the BAM file, keeping the network load for NFS storage and time-expensive IO on a high rate. Table 5 shows the sorting runtime performance of Samtools for the dataset IV run with different amount of RAM available on a single HPC node. The large uncertainty for small amounts of RAM we believe is due to other users' load on the cluster network. As a result, a large amount of RAM per node is needed in order to use Samtools effectively, while Hadoop accomplishes SNP calling on much more economic hardware in linear time with just 12GB RAM per

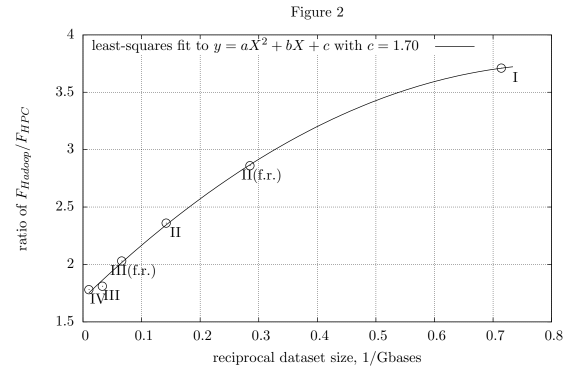


Figure 2: The ratio of the F_{Hadoop}/F_{HPC} as a function of a reciprocal dataset size in Gigabases. Calculations were carried out for 56 cores Hadoop and 16 cores HPC cluster correspondingly. The points are fit to a quadratic least-squares curve, which makes it possible to predict *infinite* dataset size. Datasets are labeled with roman numerals.

virtual node.

3.5 Comparing the network communication efficiency for the Hadoop and HPC approaches

The network communication model for Hadoop has a major difference from the usual HPC cluster network architecture (*n*-tier tree) with NAS or NFS attached storages. The effective bandwidth of the Hadoop network increases with the cluster size [13], opposite to that of the HPC network where cluster growth results in network saturation and performance depletion. To study this phenomena, we compared HPC and Hadoop network communication costs depending on the number of nodes involved for a fixed dataset size (58 Gb, dataset IV).

Due to the trivially parallelisability of the alignment process – the read-pairs are independent of each other, and can be aligned independently – one could try to involve more computational resources, e.g., split the initial data into chunks to process them independently. Reducing the size of each data chunk reduces the aligner job, $T_{mapping}$, but at the same time, the more chunks almost simultaneously have to be sent over the network, potentially causing traffic jams, and therefore increasing the communication costs, T_{comm} .

Table 5: Timings (in minutes) for the sort routine of the Samtools package for the dataset IV (58Gb BAM file). The measurements are done on a single HPC node with p=16 cores. Averaging is done over 5 independent simulations. The amount of RAM is given for all 16 cores.

RAM, GB | 8 | 16 | 32 | 64
timing, minutes |
$$208 \pm 50$$
 | 172 ± 13 | 136 ± 12 | 87 ± 11

There are several program packages for short read alignment with MPI support [7, 9], and almost linear scaling up to 32 nodes for pair-ended reads has been reported [22]. However, e.g., the Pmap package functions poorly for datasets larger than 20 Gbases. To circumvent this limitation we implemented a highly optimized Bash script making use of standard Unix utilities to use the HPC cluster network as efficiently as possible [?], and compared the network performance with the standard Hadoop HDFS approach. We separated the mapping time, $T_{mapping}$, and the communication time, T_{comm} , and plotted $T_{mapping}/T_{comm}$ as a function of the reciprocal number of nodes 1/N (Figure 3). This measure is applicable to both HPC and Hadoop, however, T_{comm} has a different explanation. For Hadoop, the short reads in FASTQ format have to be preprocessed (involving node communication T_{comm}) to be able to run in MR-fashion, while the data locality will be automatically achieved during the data ingestion into the HDFS. We rewrote the code for the preprocessing stage for Crossbow to make it suitable for MR-style parallelisation. For the HPC approach, T_{comm} involves the chunks that are sent from the sequence delivery location to the local node scratch disks where the actual mapping happens, and the sending of the aligned SAM files back to the delivery location over the network.

Hadoop results (filled circles in Figure 3): One can see that the ratio $T_{mapping}/T_{comm}$ reveals very weak dependency in a wide range of number of nodes N: from 4 up to 40. It is known that Bowtie provides almost linear scaling between mapping time and dataset chunk size D: $T_{mapping} \propto D \propto 1/N$, see [2], and Figure 1. Since the ratio $T_{mapping}/T_{comm}$ is approximately constant, one can conclude that $T_{comm} \propto 1/N$, meaning that the more nodes involved, the faster the communication is in the preprocessing stage.

HPC results (open circles in Figure 3): The data from the delivery location is split into chunks in parallel, which are simultaneously pushed to the local scratch disks of the nodes allocated by SLURM. One can see two distinct linear stretches. One stretch is for the range from 4 to about 12 nodes, and the another is from 12 up to 60. The former (horizontal stretch) is explained as for Hadoop – the more nodes involved, the faster the chunks are being distributed. The latter stretch with the positive slope could be explained as follows: In the region of about 12 nodes the network becomes saturated and unable to pass more data in a time unit, while the mapping time is still proportional to the chunk size: $T_{comm} \approx \text{const}$, $T_{mapping} \propto D \propto 1/N \rightarrow T_{mapping}/T_{comm} \propto 1/N$, i.e., a linear dependency, which can be observed in the plot. The transition area between the two modes is based on the fact that each hard disk drive on the local node can write the data at a speed of about 1Gbit/sec. Ten nodes will consume the data with the rate of 10Gbit/sec, which is the limiting speed for the standard 10Gbit Ethernet cable connecting the cluster's rack with the switch. The nodes are being allocated on the same rack, which is the default

Ratios of the short reads mapping time per node to the communication time

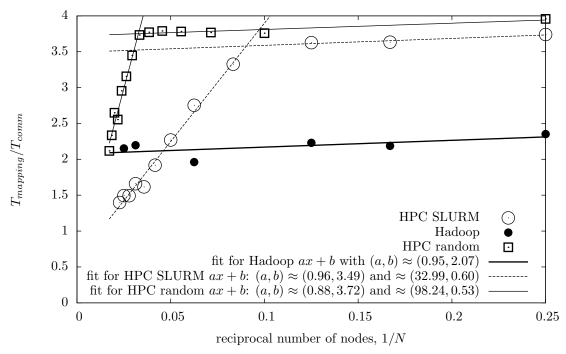


Figure 3: Ratios of the mapping time $T_{mapping}$ to the communication costs T_{comm} for HPC and Hadoop clusters for Dataset IV as a function of reciprocal cluster size 1/N. Two HPC scenarios are shown as "HPC SLURM" and "HPC random", which correspond to standard SLURM behaviour and a modified one where nodes are being allocated from random racks. Linear fit done with the least-squares method. Figure legend too complex. Y-axis label inconsistant with previous figure.

SLURM behaviour.

Scalability can be improved by overriding the default behaviour of SLURM and allocating the nodes not from the same rack, but randomly from all available racks ("HPC random", open squares in Figure 3). Allocating the nodes on random racks allows one to engage more nodes without network saturation. For our cluster we could go up to 30-35 nodes with perfect linear scaling. For the most resources used (58 nodes) the deviation from a linear speedup is $\approx 7\%$ i.e. 5.50 minutes against the ideal 5.14, see Table 4. The threshold number of nodes in this strategy (≈ 35) is because of the uplink cable with the throughput of 50Gbit/sec being saturated. The proposed HPC strategies aimed at getting the maximum performance from the storage resources show that while even properly adjusted and tuned, the HPC approaches suffer from the network saturation at higher number of nodes.

HDFS on the other hand maintains data locality, reducing communication and data transfer and hence lead to better scalability. Our Hadoop cluster has no more free nodes to continue to investigate the scaling as in plot at Figure 3, but we do not expect any significant deviations, since the observed behaviour is a generic for Hadoop with HDFS. need ref for this!

Table 6: Timings for mapping and the ratio $T_{mapping}/T_{comm}$ for HPC and Hadoop clusters for Dataset IV. For the "HPC random" approach, data chunks have to be copied to the local scratch disks first and the alignments (SAM files) copied back while Hadoop keeps all the data inside HDFS and hence does not need data staging. Hadoop however needs to preprocess reads before the actual alignment stage in order to be able to operate in MR manner resulting in what we term "communication costs". Note that each HPC node has 16 cores, while each Hadoop node has 7 (one core is dedicated to run the virtual machine).

	Hadoop		HPC random			
Number of nodes	Mapping time,	$\frac{T_{mapping}}{T_{comm}}$	Number of nodes	Mapping time,	$\frac{T_{mapping}}{T_{comm}}$	
(cores)	minutes	- comm	(cores)	minutes	- comm	
4(28)	293.5	2.33	4(64)	74.4	3.89	
6(42)	189.8	2.19	10(160)	32.4	3.76	
8(56)	136.0	2.23	14(224)	22.7	3.77	
16(112)	70.3	1.96	18(288)	17.9	3.78	
32(224)	39.3	2.20	22(352)	14.5	3.79	
40(280)	32.5	2.15	26(416)	12.3	3.77	
			30(480)	10.7	3.73	
			34(544)	9.5	3.45	
			38(608)	8.5	3.16	
			42(672)	7.6	2.96	
			46(736)	7.0	2.55	
			50(800)	6.4	2.65	
			54(864)	5.9	2.34	
			58(928)	5.5	2.12	

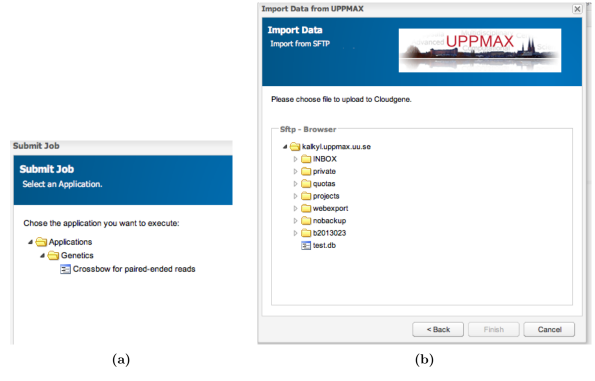


Figure 4: An example of a job setup with the graphical Hadoop front-end Cloudgene, providing a smooth user experience even for novice users. a) Pipeline selection - in our case containing the Crossbow pipeline. b) The UPPMAX-adapted functionality to browsing and import data from the user's home folder in the shared file system.

3.6 Usability aspects

A popular way to construct and execute bioinformatic pipelines on HPC resources is via the Galaxy[23] platform, which provides a Web-based graphical user interface (GUI) to bioinformatic programs, simplifying the experience for the end user. An alternative for Hadoop is Cloudgene[24], which is a light-weight and flexible Web-based solution for both public and private clouds. We implemented our Hadoop-pipeline in Cloudgene and extended the platform with functions to import data from the central file system at UPPMAX into HDFS on the private cloud (Figure 4). For our particular task in DNA sequencing, Cloudgene makes it easy to select data and parameters for its execution. Most of the data managing work is done automatically and the results can be downloaded to the client machine. The modular structure allows modification of the source code to adapt to the existing computing center architecture. For example, UPPMAX users can import their data from the sequencing platform directly to the Hadoop cluster by pressing a button and entering the credentials, being at the same time sure that their sensitive data is kept privately.

4 Discussions/Conclusions

In this report we have described two approaches for high-performance analysis of DNA sequencing data; one based on regular HPC using a batch system and one based on Hadoop. We show that Hadoop installed on a private cloud is an appealing solution both on terms of runtime performance and also in terms of usability. A key result is the data size where Hadoop is favorable to regular HPC batch systems, and in order to compare them we developed highly optimized pipelines for both scenarios. We found that dataset sizes larger than 100 Gbases is where Hadoop excels over HPC, and also that Hadoop shows almost linear scalability for this problem. Extrapolation to infinite dataset size reveals however that HPC provides the results faster, given the same amount of resources as Hadoop on a private cloud.

We modified the preprocessing stage of the Crossbow software to enable it to make use of all nodes in the Hadoop system, end used a highly optimized Bash script to compare the scaling relations between the ratio of the mapping time to the communication time as a function of the reciprocal number of nodes on an HPC system. Our results show that the calculations on Hadoop with HDFS scales better than the network attached parallel storage commonly used in the HPC centers. In addition we show how performance of the HPC approach can be improved by redefining the queueing system's default behaviour. This is however not of practical use in an HPC center since the network bandwidth is shared among multiple users. Finally, we demonstrate that an existing and extensible publically available webbased GUI (Cloudgene) provides an easy way to execute bioinformatics analysis on Hadoop for those who are less experienced with Linux scripting.

5 Acknowledgements

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6 Supplementary material

6.1 Datasets

The datasets used in the paper are publicly available at:

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I: http://1001genomes.org/data/software/shoremap/shoremap_2.0\/data/reads/Schneeberger.2009/Schneeberger.2009.single_end.gz
II: http://1001genomes.org/data/software/shoremap/shoremap_2.0/data/
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reads/Galvao.2012/Galvao.2012.reads1.fq.gz,http://1001genomes.org/data/software/shoremap/shoremap_2.0/data/reads/Galvao.2012/Galvao.2012.reads2.fq.gz

III: ftp://ftp-trace.ncbi.nlm.nih.gov/sra/sra-instant/reads/ByRun/sra/
SRR/SRR611/SRR611084//SRR611084.sra,ftp://ftp-trace.ncbi.nlm.nih.gov/sra/
sra-instant/reads/ByRun/sra/SRR/SRR611/SRR611085//SRR611085.sra

IV: artificial pair-ended dataset for A.thaliana created with the wgsim program from the Samtools package.

V: http://www.ncbi.nlm.nih.gov/sra/SRX148888

6.2 Reference genomes

- TAIR10 for datasets II-IV ftp://ftp.arabidopsis.org/home/tair/Sequences/whole_chromosomes/*.fas
- TAIR8 for dataset I ftp://ftp.arabidopsis.org/home/tair/Genes/TAIR8_genome_release/
- H.sapiens, NCBI v37 ftp://ftp.ccb.jhu.edu/pub/data/bowtie_indexes/ h_sapiens_37_asm.ebwt.zip

6.3 Description of computational facilities

- 1. HPC: Multinode short read mapping was performed on the Milou cluster [25], equipped with dual 8-core Intel Xeon E5-2660, (2.2 GHz, 2MB L2 cache, 20 MB L3 cache), 128 GB of RAM, Infiniband node-to-node network connection, and 10 Gbit/s uplink.
- 2. Storage: Gulo[26] is a custom built Lustre 2.4 system using 8 HP nodes with MDS600 storage boxes and an additional node for metadata handling. In total, it provides roughly 1 PB of storage and is accessed with Lustre's own protocol. It supports data striping over multiple nodes and disk targets and can give a theoretical single file read performance of up to 80 Gbit per second.
- 3. Our Hadoop test platform was deployed on a private cloud at UPPMAX using the OpenNebula [18] cloud management system. Each node in this deployment was equipped with dual 4-core Intel Xeon 5420 (2.50 GHz; 12 MB L2 cache), 16 GB RAM, one 1 TB SATA disk and Gigabit Ethernet. The cluster was set up with Cloudera Hadoop Distribution version 2.0.0-cdh4.4.0 [19]. Note that the physical hardware provided less RAM than desired. Each VM node has 7 cores where each can use less than 2 GB of RAM, which is little for Hadoop. We expect much better performance with twice as much memory.

6.4 Bash script for the preprocessing stage

```
pbzip2 -dc $file1 | paste - - - - -d'\t' | cut -f1,2,4 | paste - -d' '
<(pbzip2 -dc $file2 | paste - - - - -d'\t' | cut -f2,4) | pbzip2 -cz >
$fileOut
```

Journal abbreviations in references are inconsistant. The journal name for ref 2 is wrong. Capitalization of refs 4, 14, 16, 18, 19, 25, 27, etc are wrong. All refs should be checked for accuracy and style consistancy. Many or all footnotes should be changed to endnotes (references).

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