

REVIEW ARTICLE

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# Data compression for sequencing data

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## Abstract

Post-Sanger sequencing methods produce tons of data, and there is a general agreement that the challenge to store and process them must be addressed with data compression. In this review we first answer the question “why compression” in a quantitative manner. Then we also answer the questions “what” and “how”, by sketching the fundamental compression ideas, describing the main sequencing data types and formats, and comparing the specialized compression algorithms and tools. Finally, we go back to the question “why compression” and give other, perhaps surprising answers, demonstrating the pervasiveness of data compression techniques in computational biology.

## Background

In the first decade of the century, the cost of sequencing a single human genome fell from about 30 million to about 10 thousand dollars (here and later we mean U.S. dollars). The second generation sequencing platforms by 454 Life Sciences, Illumina, and Applied Biosystems cost less than one million dollars [1] and are available in many institutes. The promised third generation (3G) technology (including Ion Torrent Systems, Oxford Nanopore Technologies, and Pacific Biosciences equipment) should be even cheaper, which rapidly moves us closer to the day of personalized medicine available to the masses.

In mid-2013, the world-biggest sequencing institute, Beijing Genomics Institute, used 188 sequencers, of which 139 were the top-of-the-line Illumina HiSeq 2000 and 2500 sequencing machines. Their total theoretical throughput is over 1.2 Pbases per year, which is equivalent to about 3 PB of raw sequencing read files. Including the additional output space for mapping to the reference genomes, the total amount of necessary storage is on the order of 10 PB per year. The statistics of high-throughput sequencers in the world (<http://omicsmaps.com>) show that the storage necessary for the instruments’ output is in the range 50–100 PB per year. Kahn [2] presents the genomic data growth until 2010, pointing out that the progress in computer hardware lags behind.

Interestingly, recently the Million Veteran Program (MVP), led by the US Department of Veterans Affairs, was announced. With at least 30-fold coverage (100 bp reads) the number of reads per genome sample will be about 1 billion [3]. This means about 250 PB of raw data (in FASTQ format) in total, when the sequencing program is finished (the enrollment of volunteers is expected to last 5 to 7 years).

Those numbers are very large, but we need to remember that they refer to July of 2013. As can be seen in Figure 1, the cost of sequencing a single base has been halving roughly every 8 months in 2008–2013, while the cost of hard disk space has been halving every 25 months in 2004–2013. Even if the most recent NHGRI data suggest some stagnation (see also the comment [4]), it may be a temporary slowdown, as 3G instruments are becoming available (PacBio RS and Heliscope).

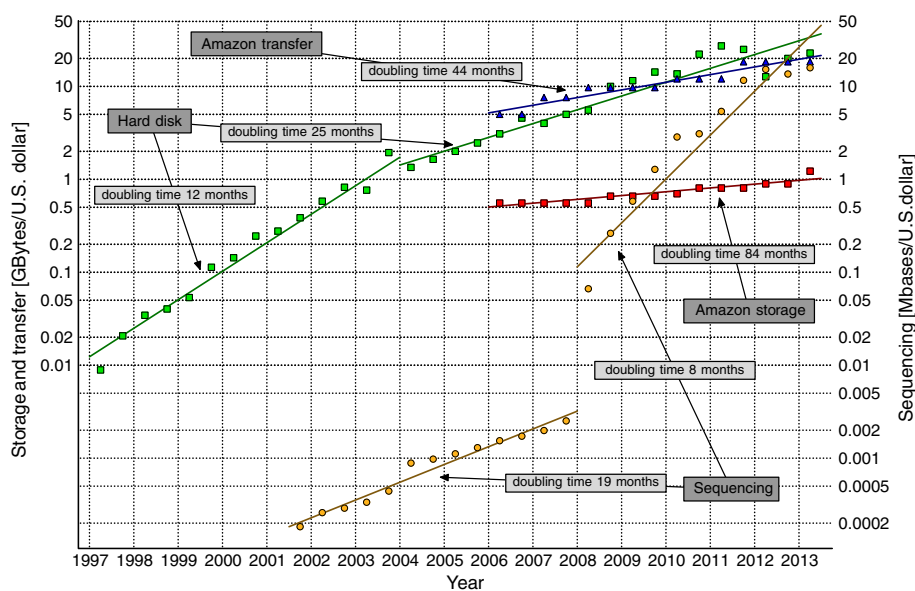
On the other hand, the prices of low-end hard disks are rather misleading, since the amount of data necessary in sequencing projects is so huge that data centers are much better places to store the files. The growing popularity of cloud storage can be attributed to several reasons. Generally, data management may be cheaper if run by IT professionals (not always available in smaller centers, like hospitals), centralization may reduce data replication costs, and access to really huge repositories (at least of terabyte scale) is easier or even only possible with large arrays of disks. The disk drives in large storage arrays are usually enterprise-class ones of increased reliability and better performance, so they are several times more expensive than standard (SATA) HD drives. Taking these factors together we should not be surprised that storing a file in a

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**Figure 1 Trends in storage, transfer, and sequencing costs.** The historic costs of low-end hard disk drives were taken from <http://www.jcmit.com/diskprice.htm>. They had been halving every 12 months in the 1990s and around 2000–2004. Then, the doubling time lengthened suddenly, to about 25 months. The real costs of sequencing, taken from the NHGRI Web page [5], reflect not only reagent costs like some studies show, but also include labor, administration, amortization of sequencing instruments, submission of data to a public database, etc. The significant change in sequencing costs around 2008 was caused by the popularization of the second generation technologies. The prices of the Amazon storage and transfer reflect the real market offers from the top data centers. It is interesting that the storage costs at data centers drop very slowly, mainly because the costs of blank hard disks are only a part of the total costs of maintenance. The curves were not corrected for inflation.

large data center may cost about 5–10 times more than on a plain HD; this is the price we pay for ubiquitous access to reliable data collections.

What is more, these files must be transferred over the Internet, which is not free. One of the largest data centers and cloud computing centers are Amazon S3 and Amazon EC2. Their charges for storage has been halving every 84 months and the charges for transfer has been halving every 44 months in 2006–2013. In January 2013 the real cost of sequencing human-size genome (according to NHGRI data) was about 5,700 dollars, while the cost of one-year storage at Amazon S3 and 15 downloads of raw reads and mapping results (225 GB of reads with 30-fold coverage and 500 GB of mapped data) was close to 1,500 dollars<sup>a</sup>.

The trends show clearly that the costs of storage and transfer will become comparable to the costs of sequencing soon, and the IT costs will be a significant obstacle for personalized medicine, if we do not face this problem seriously.

When choosing the compression algorithm to apply (if any), one should consider the specific use of the data. Two, in a sense extreme, scenarios are: (i) the data are to be transferred over a network (and then decompressed), (ii) the data are to be accessed in real-time, either in a sequential or random fashion. Let us take a closer look at both. In the case (i), a reasonable cost measure is the time to

compress the given data (preferably using some “standard” computer), transfer it, and decompress on the receiver side. Concerning sequencing reads (FASTQ), it is possible to obtain (at least) 4.5-fold compression at about 30 MB/s both compression and decompression speed, on a single CPU core. Let us assume a link speed of 50 Mbit/s and that the input files have 450 GB in total. Without any compression the files will be transferred in exactly 20 hours. On the other hand, the compression, transfer and decompression times sum up to about 10.5 hours. Even if a faster connection is available (when the gain diminishes), we note that the compression and decompression may also be sped up (using threads or separate processes to utilize multiple CPU cores, and also by overlapping the compression, transfer and decompression phases). We point out that the ratio of 4.5, used in this example, is rather modest; significantly more is possible with lossy compression and also for some other biological data types (like genome collections).

Let us now estimate the monetary savings for transferring (downloading) these files from Amazon S3. The transfer charge depends on the volume of data, but let us assume 9 cents/GB. Without compression, we will pay 40.5 dollars for downloading these 450 GB of data, but only 9 dollars if compression is applied.

The case (ii) is somewhat different, as the choice of the compression method is rather constrained. To our knowledge, the most successful, in terms of efficiency

and flexibility, storage system for biosequences was presented by Steinbiss and Kurtz [6]. Their GtEncseq solution accepts several common sequence formats (FASTA, GenBank, EMBL, FASTQ), compresses the input, provides fast random access to FASTA data, stores metadata, etc. Accessing single bases from random locations is not much slower than from the uncompressed equivalent representation, with over 3 million queries per second. In case of FASTQ, however, extracting a single read from an arbitrary location is not that fast, as it takes, e.g., more than 100  $\mu$ s, which is less than the access time of a hard disk drive, but not necessarily faster than the SSD. A space-time tradeoff exists, so this time may be reduced, but consequently compression ratio drops quite severely. Anyway, if only sequential access to FASTQ is needed (which is often the case), GtEncseq may be a good choice.

As new large-scale sequencing projects are frequently announced (the mentioned MVP project is probably the most ambitious at the moment), the storage concerns are of high priority. It is no wonder then that many ideas of selective storage (or lossy compression) are discussed in the community (cf. [7,8]). While we agree that some forms of lossy compression seem to be a necessity, the radical approach of discarding old data (in the hope of reproducing them later, when needed) raises major methodological doubts, as far as research applications are considered. More precisely, discarding raw data is hazardous, since their reproduction later will not be exact, due to inherent “randomness” in sequencing process, even if the same hardware and possibly identical procedure are used. This clashes with one of the main principles of the scientific method, which is reproducibility.

The area of data compression techniques in computational biology has been surveyed by Giancarlo *et al.* [9,10], with more focus on the theory and data compression applications in sequence analysis than storage and indexing of data from high-throughput technologies. One aspect of data compression in genomics, index structures for sequencing data, is thoroughly discussed by Vyverman *et al.* [11]. We hope our review complements earlier efforts by paying attention to versatile applications of data compression in the age of data flood in bioinformatics.

The paper is organized as follows. The next section contains a brief introduction to widely used data compression algorithms. The section “Sequencing data” outlines the popular compression techniques and file formats to store various bioinformatics data: base calls annotated with quality scores, genome alignment data, single and multiple genome data. Making data smaller is not only for reducing their space and facilitating their distribution, as it is reflected in the name of the next section, “Beyond storage and transfer”. Here, applications of data

compression ideas in indexing, read alignment and other vital problems are discussed. The last section concludes.

#### Data compression in brief

Compression techniques are the traditional means of handling huge data. Those methods reduce the space for storage and speed up the data circulation (e.g., among research institutes). With its origins yet in the 19th century and first theoretical works just after the WW2, nowadays data compression is used almost everywhere as the amount of stored and transmitted data is huge. There are several major concepts that are used in compression programs, but the two mentioned below are the most important in bioinformatics. Here we give only a very short description of them, with more details in the Additional file 1 or the monograph [12].

The Huffman coding [12,13], invented in 1952, is a statistical method, which assigns a sequence of bits (a *codeword*) to each alphabet symbol. The codewords are of different length, and in accordance to the golden rule of data compression, rarer symbols are represented by longer codewords. The given sequence is then encoded by replacing each symbol with its corresponding codeword. What makes Huffman coding important is its optimality, i.e., no other code leads to a shorter encoded sequence.

In 1977–78 Ziv and Lempel [12,14] invented dictionary methods. They process the sequence from left to right and encode possibly long repetitions of consecutive symbols as references to the already compressed part of data. Such an approach allows for higher compression ratios than sole Huffman coding as it looks for another type of redundancy, common not only in natural language (e.g., repeating word phrases), but also in multiple genome sequences or overlapping sequencing reads. Even better results are possible with combining dictionary methods and Huffman coding. The very popular gzip program serves as a successful example.

The Burrows–Wheeler transform (BWT) [12,15] is a more recent compression idea, that has become highly popular in bioinformatics. The pure BWT is not a compression method, it only permutes the input sequence, but it can be used to construct highly efficient compressors (bzip2, combining the BWT with Huffman coding and other techniques, is a well-known representative). This transform is also a basis of some sequence indexing techniques that are used to search a genome in many tools (more on this will be covered in the “Beyond ...” section). The key idea of BWT is to permute the input sequence in such a way that symbols are grouped by their neighborhood, i.e., the symbols that are followed by the same symbols are close after the permutation, even if they were far in the original sequence (see also the Additional file 1 for a BWT example).

## Sequencing data

### Raw sequencing data

The raw reads from sequencing are usually stored as records in textual FASTQ format. Each record is composed of a read ID, base calls, and quality scores for all base calls [16]. These files are often compressed with gzip, to obtain about 3-fold size reduction. While significant, such a gain is not quite satisfactory. A better choice is to use a FASTQ-dedicated compressor, e.g., DSRC [17], which shrinks the data about 5 times. The algorithm finds overlaps in the base calls from the reads placed at very long distances in the file (in Ziv–Lempel fashion), as well as takes care of various ID format conventions. It also uses multiple statistical models to better compress quality values by the Huffman coding. In particular, quality scores in different read positions are compressed in different models (i.e., based on different statistics) because it is well-known that the quality deteriorates towards the read end, hence this design improves the prediction of the scores. DSRC is quite fast, as it compresses at 30–40 MB/s speed, and handles alphabets of size beyond 5 (IUPAC ambiguity codes) as well as variable-length reads.

Most newer proposals [18,19] focus mainly on compression ratio and either the speeds are of secondary importance or are not examined at all. For example, the solution by Bholá *et al.* [18] follows essentially the same direction as DSRC, but handles also approximate and palindromic repeats. The byproducts are compressed with an adaptive arithmetic coder [12]. While the reported compression ratios are usually higher than the DSRC's by a few percent, it is unclear if the solution can be scalable. Remarkably, doubts are expressed by the authors themselves.

A more radical attempt to improve compression ratio is to group the reads with suffix-prefix overlaps close together. Unfortunately, the algorithms following this line [20,21] are not fully functional tools, e.g., they ignore the read IDs and quality scores, and achieve processing speeds on the order of 1 MB/s or less. A more interesting solution along these lines, SCALCE [22], is a FASTQ preprocessor helping to improve the compression ratio with gzip twice.

As argued earlier, the rapid growth of data from sequencing experiments demands even better compression ratios, and switching to a lossy mode seems to be the only chance for a breakthrough [23]. The natural candidates for lossy compression are quality scores. It has been shown [22,24–26] that rounding the quality scores to a few values (instead of about 60) can be acceptable. E.g., the fraction of discrepant SNPs grows slowly with diminishing number of quality scores in Illumina's CASAVA package [27] while the benefit in compression is clear<sup>b</sup>. On the other hand, some (re)assemblers ignore these data [23], so if the reads are to be processed by such tools, the quality scores can be removed. Another option is to use the

scores to prefilter the reads and discard the ones with unacceptably low quality.

SeqDB [28] is a notable exception from the shown tendency. Here the focus was put on compression and decompression speed, not compression ratio. On a machine with 12 CPU cores SeqDB reaches the speeds of about 500–600 MB/s ([28], Section 4.4), but the compression ratio is at best at gzip's level.

Recently, two very interesting algorithms were presented. One of them is Quip [29], which uses higher-order modelling [12] with arithmetic coding in its more "traditional" mode, and also an assembly-based approach in its stronger mode. The idea is to form contigs from the first (by default 2.5 million) reads, which then are used as a reference for the following reads. The compression improvement due to this idea is moderate however, so the standard mode (faster and using half of the memory) seems to be more practical, with compression significantly better than DSRC and 10 MB/s to 20 MB/s compression/decompression speed.

In [30] two FASTQ compressors were presented. Now we briefly describe the more interesting of them, Fqzcomp. It achieves better compression ratios thanks to a carefully chosen context mixing model<sup>c</sup> and other original ideas. Fqzcomp belongs also to faster solutions, (partly) due to multi-threading. In the same paper, yet another strong FASTQ compressor was tested, SeqSqueeze1 (described only in [30]). This algorithm, also based on context mixing, is sometimes even better than Fqzcomp in respect of compression ratio, but its compression and decompression speed is less than 1 MB/s.

In Table 1 we compare the existing tools for compressing sequencing data. They belong to four different kinds described in the following four subsections. We did our best not to miss any relevant tool for which the sources or (at least) binary executables for a popular operating systems were available. There are several reasons why we do not present experimental (comparative) tests of the presented tools. They are, in the order of decreasing importance:

- (i) limitations of many tools (e.g., accepting sequences with only the ACGT or only ACGTN alphabet, support for fixed-width reads only, assumptions on the ID format in FASTQ files, restriction to only selected fields of the SAM format);
- (ii) significant problems with running some of the existing tools (which we have experienced in earlier work, on FASTQ and genome collection compression);
- (iii) not quite compatible outputs (e.g., DSRC for FASTQ supports random access, which cannot be turned off, while many others do not, hence comparing compression ratios of these tools cannot be fully fair);

**Table 1 Summary of the most important compressors of sequencing data**

Software name	Implementation availability src code / binaries / libs	Website	Lossless / lossy	Ambig. codes	Var. length reads	Speed of compr./decompr.	Ratio	Random access	Methods	Remarks
<b>Compressors of raw sequencing data</b>										
gzip	C++ / many / many	www.gzip.org	yes / no	yes	yes	moderate / very high	low	no	LZ, Huf	
bzip2	C / many / many	www.bzip.org	yes / no	yes	yes	low / high	low	no	BWT, Huf	
7zip	C, C++ / many / many	www.7-zip.org	yes / no	yes	yes	low / very high	moderate	no	LZ, AC	
BWT-SAP [21]	C++ / — / C++	github.com/BEETL/BEETL	yes / no	yes	no	low / low	moderate	no	BWT, PPM	FASTA only
DSRC [17]	C++ / Lin, Win / C++, Pyt	sun.aei.polsl.pl/dsrc	yes / no	yes	yes	high / high	moderate	yes	LZ, Huf	
Fqzcomp [30]	C / Lin / —	sourceforge.net/projects/fqzcomp/	yes / yes	no	yes	high / moderate	high	no	CM	
G-SQZ [31]	C++ / Lin / —	public.tgen.org/sqz	yes / no	no	no	high / moderate	low	yes	Huf	
Kung-FQ [19]	C# / Win / —	quicksaf.sourceforge.net	yes / yes	no	no	moderate / moderate	moderate	no	AC, LZ, RLE	
Quip [29]	C / — / —	cs.washington.edu/homes/dcjones/quip	yes / no	no	no	high / high	high	no	M. models, AC	
ReCoil [20]	C++ / — / C++	github.com/BEETL/BEETL	yes / no	no	no	very low / high	moderate	no	BWT, PPM	FASTA only
SCALCE+gzip [22]	C++ / — / —	scalce.sourceforge.net	yes / yes	no	no	moderate / high	moderate	no	AC, LZ, Huf	
Seq-DB [28]	C++ / — / —	https://bitbucket.org/mhowison/seqdb	yes / yes	no	no	very high / very high	low	yes	AC, LZ, RLE	
SeqSqueeze1 [30]	C / Lin / —	sourceforge.net/p/ieetaseqsqueeze/wiki/Home/	yes / no	no	yes	very low / ver low	high	no	CM	
<b>Compressors of reference genome alignment data</b>										
gzip	C++ / many / many	www.gzip.org	yes / no	yes	N/A	low / very high	low	no	LZ, Huf	
bzip2	C / many / many	www.bzip.org	yes / no	yes	N/A	low / high	low	no	BWT, Huf	
7z	C, C++ / many / many	www.7-zip.org	yes / no	yes	N/A	low / very high	moderate	no	LZ, AC	
BAM [32]	C++ / many / many	samtools.sourceforge.net	yes / no	yes	N/A	moderate / high	moderate	yes	LZ, Huf	
CRAM [33]	Java / many / Java	www.ebi.ac.uk/ena/about/cram_toolkit	yes / yes	yes	N/A	moderate / moderate	moderate	yes	Huf, Gol, diff.	
Quip [29]	C / — / —	cs.washington.edu/homes/dcjones/quip	yes / no	no	N/A	high / high	high	no	M. models, AC	
SAMZIP+rar [34]	C / — / —	www.plosone.org/article/info:doi/10.1371/journal.pone.0028251	yes / no	yes	N/A	moderate / high	moderate	no	RLE, LZ, Huf	

**Table 1 Summary of the most important compressors of sequencing data (Continued)**

<b>Compressors of single genome sequences</b>									
gzip	C++ / many / many	www.gzip.org	yes / no	yes	N/A	moderate / very high	low	no	LZ, Huf
bzip2	C / many / many	www.bzip.org	yes / no	yes	N/A	low / high	low	no	BWT, Huf
7z	C, C++ / many / many	www.7-zip.org	yes / no	yes	N/A	low / very high	moderate	no	LZ, AC
dna3 [35]	C / - / -	people.unipmn.it/manzini/dnacorpus/	yes / no	no	N/A	low / low	moderate	no	LZ, PPM
FCM-M [36]	C / - / -	ftp://www.ieeta.pt/~ap/codecs/	yes / no	no	N/A	very low / very low	moderate	no	M. models
XM [37]	Java / many / Java	ftp.infotech.monash.edu.au/software/DNAcompress-XM	yes / no	yes	N/A	very low / very low	moderate	no	M. models, AC
<b>Compressors of genome collections</b>									
gzip	C++ / many / many	www.gzip.org	yes / no	yes	N/A	low / very high	very low	no	LZ, Huf
bzip2	C / many / many	www.bzip.org	yes / no	yes	N/A	low / high	very low	no	BWT, Huf
7z	C, C++ / many / many	www.7-zip.org	yes / no	yes	N/A	low / very high	high	no	LZ, AC
ABRC [38]	C++ / Lin, Win / C++	www2.informatik.hu-berlin.de/~wandelt/blockcompression/	yes / no	yes	N/A	high / very high	very high	yes	LZ, Huf
GDC [39]	C++ / Lin, Win / C++	sun.aei.polsl.pl/gdc	yes / no	yes	N/A	high / very high	very high	yes	LZ, Huf
GReEn [40]	C / - / -	ftp://ftp.ieeta.pt/~ap/codecs/	yes / no	yes	N/A	high / high	high	no	M. models, AC
GRS [41]	C / Lin / -	gmdd.shgmo.org/Computational-Biology/GRS/	yes / no	yes	N/A	moderate / low	high	no	LCS, Huf
RLZ [42]	C++ / - / -	www.genomics.csse.unimelb.edu.au/product-rlz.php	yes / no	yes	N/A	moderate / very high	high	no	LZ, Gol

Abbreviations used in the table: src—source codes, libs—libraries, Lin—Linux, Win—Windows, Pyt—Python, exe—binary executables, AC—arithmetic coding (a statistical coding method [12]), CM—context mixing for arithmetic coding [12], diff—differential coding (paradigm: store only changes between sequences), Gol—Golomb (a statistical coding method [12]), Huf—Huffman, LCS—longest common subsequence (a measure of similarity of sequences [43]), LZ—an algorithm from Ziv–Lempel family, M. models—Markov models [12], PPM—prediction by partial matching (an efficient general-purpose compressor [12]). “Ambig. codes” means the ability to compress DNA symbols other than {A, C, G, T, N}. “chr-ordered” for 7z and genome collections means that the input (human) genomes were split into chromosomes and ordered according to them before the actual compression. In this way several chromosomes fit the 7z LZ-buffer which is beneficial for the compression.

- (iv) different targets (by design, some of the tools are supposed to be run on a commodity PC, while others require powerful servers, or never were tested on mammalian-size data, since it would take days or even weeks. For example, many top single genome sequence compressors work at the rate of several Kbase/s only, on a standard computer).

### **Reference genome alignment data**

The reads are usually assembled or reassembled. *De novo* assembling is the most challenging, but resequencing, in which the reads are aligned to some reference genome, is much cheaper and thus widely used.

The results of mapping the reads onto the reference genome are usually stored in the SAM/BAM format [32]. SAM files augment the reads data with mapping quality and several other fields. BAM is a gzip-like compressed binary equivalent of textual SAM and is about 3–4 times smaller. Due to the additional data SAMs are more than twice as large as FASTQ files.

The reads in SAM files are mapped to a known reference genome and the differences between the reads and the reference sequence, resulting from variation and sequencing errors, are small. Thus, it is efficient to represent the base calls of a read as the mapping coordinate and the differences. These reads are usually ordered by the mapping coordinate and thus the coordinates can be stored in a differential manner, which results in a sequence of small and thus well compressible numbers. The oldest scheme [44] for compressing mapping data with the described idea cannot however be considered mature, since quality scores are ignored and there is no support for unaligned (i.e., those that failed to map onto a reference) reads.

Fritz *et al.* [33] handle both aligned and unaligned reads. The aligned reads are stored basically as described above together with the quality scores. To obtain better compression ratios, the authors advocate using a lossy mode and refrain from storing some quality scores, e.g., the ones related to perfectly matched positions. To compress the unaligned reads (usually 10–40% of raw reads) better, they propose to build some artificial reference sequences. To this end, unmapped reads from many similar experiments are processed by an assembler to obtain contigs, built only for the compression process. Finally, the remaining sequences are matched to the bacterial and viral databases. Some of the byproducts in the algorithm are encoded with the Huffman (or other) codes. Most of the described techniques, excluding artificial reference as well as bacterial and viral sequences, are implemented in the CRAM compressor. In a highly lossy setting, it can produce archives smaller by an order of magnitude than corresponding BAM files. Similar approaches for compressing reads by mapping them onto a reference

genome are used in SlimGene [25] and SAMZIP [34] algorithms.

Generally the use of a reference sequence can help a lot for the compression ratio but we should remember that a reference might not be available in some cases, e.g., for metagenomic datasets or for organisms with high polymorphism [20].

The most recent SAM compressor [45], apart from highly configurable lossy compression settings, introduces a novel idea to exploit common features of reads mapped to the same genomic positions. Quip [29], published slightly earlier, is not as flexible as CRAM and works only losslessly. However, if aligned reads in the SAM or BAM format and a reference sequence are given, it wins with CRAM in compression ratio and needs less memory to operate.

The tabix program [46] is a more general solution, popular in sequencing centers. It was designed to allow fast random access to compressed (gzip-like) textual files, in which the data are stored in rows containing tab-delimited values. The basic idea is to sort the input rows according to the sequence name and coordinate. Then the file is split into a series of blocks of maximum size of 64 KB. These blocks are compressed. Finally, an index to support random access queries is built.

### **Single genome sequences**

Raw and annotated sequencing data are nowadays the greatest challenge for storage and transfer today. Nevertheless, consensus DNA sequences (e.g., complete bacterial genomes) used to be historically the first object of compression in bioinformatics. In a sense, however, the genome sequences for a single individual are almost incompressible. If the sequence contains only the symbols A, C, G, and T, then the trivial 2 bits per symbol encoding is often more efficient than a general-purpose compressor, like gzip!

Specialized DNA compressors appeared in mid-1990s, but most solutions from the literature are impractically slow. For example, one of the strongest of them, the highly-acclaimed XM [37], can squeeze a genome up to about 5 times, but compression speed of the order of 20 KB/s on a modern machine is clearly disappointing. Some other notable compressors in this area are dna3 [35] and FCM-M [36]. The standard input format for genome sequences is FASTA, in which the file starts with a single-line description, followed by lines of sequence data.

### **Collections of genome sequences**

As said, a single genome in its compact encoding (2 bits per base) seems almost incompressible. However, large repositories with thousands of individual genomes of the same species are just behind the corner. These genomes are highly similar to each other (e.g., human genomes

have more than 99% of their content in common [47]), so a collection can be very efficiently compressed. Dictionary methods, from the LZ family, constitute the most obvious and actually most successful approach, considering high-speed decompression with moderate memory requirements. The compression phase is more demanding, both in time and space, because the repetitions (matches) in a collection of genomes are typically gigabytes apart. For these reasons, most general-purpose LZ-style compressors (e.g., gzip, rar) are useless for those data, and a few years ago the first specialized algorithms emerged.

In their seminal work, Christley *et al.* [48] compressed a single human (James Watson's) genome, but with the variation data relative to a reference genome being provided. Variation data were comprised of single nucleotide polymorphisms (SNPs) and so-called indels, i.e., insertions or deletions of multiple nucleotides. Additionally, they used a readily-available SNP database. The assumed scenario, augmented with standard compression techniques, made it possible to represent the whole human genome in about 4 Mbytes only. Recently, Pavlichin *et al.* [49] followed the lines pioneered in [48], compressing the JW genome to 2.5 MB, with very similar average results on 1092 human genomes from the 1000 Genomes Project. The introduced novelties are partly biologically-inspired, e.g., making use of the tag SNPs characterizing haplotypes. In another recent achievement, Deorowicz *et al.* [50] compressed the variant call files from 1092 diploid *H. sapiens* (1000 Genomes Project) and 775 *A. thaliana* (1001 Genomes Project) individuals, together with a variant database, squeezing the former collection to about 432 MB and the latter to 110 MB (which translates to 395 KB and 142 KB per individual, respectively).

Other research did not assume access to a knowledge base (i.e., a reference genome), hence most of them did not yet achieve the mentioned levels of compression. Most of these works [39,40,42,51] encode one of the genomes in a collection with simple means, spending about 2 bits per base, and then apply very efficient *differential* LZ-like encoding for the remaining genomes. The pioneering algorithm of this kind, RLZ [42,52], looks for LZ-matches in the reference genome (the one encoded naïvely, with 2 bits per base), and encodes their positions compactly, with reference to the previous match. This handles typical differences in genomes of the same species (short insertions or deletions, SNPs) in an efficient manner. Such an approach is successful: the related compressor, GDC [39], obtains in its strongest mode the compression ratio of 1,000 for relatively encoded genomes in a collection of 70 human individuals, with decompression speed of 150 MB/s. The key ideas in GDC include looking for long approximate matches in the whole collection and the Huffman coding.

Some of the specialized compressors (GDC, LZ-End [53]) allow for fast access to an arbitrary substring of the compressed collection. Unfortunately, this comes at a price: some loss in compression ratio (which still remains competitive, though).

Yet another differential genome compressor was presented by Wandelt and Leser [38]. Implementation-wise its original trait is searching for matches via a compressed suffix tree [54] (in blocks). This solution reaches the compression ratio of about 400 for the collection of 1000 human genomes. The match-finding speed of a parallel implementation is rather high (85 MB/s with large blocks). The real bottleneck is, however, building the compressed suffix tree.

Interestingly, also 7zip, a well-known advanced general-purpose LZ compressor, achieves quite competitive results, but for mammalian-size genomes the collections must be reordered by chromosomes, otherwise it cannot find inter-genome LZ-matches and its compression then is not much better than gzip's. This behavior can be explained by its LZ-buffer limited to 1 GB, which is less than the size of, e.g., the human genome. On the other hand, chromosomes are already small enough and several of them fit its buffer.

Currently, public repositories often store large genomes as variant databases (e.g., in VCF format). This tendency should help significantly in developing new efficient compression algorithms, since all input sequences are perfectly aligned and the (otherwise hard and resource-consuming) task of finding repetitions in data becomes almost trivial. Thus, we expect the work by Christley *et al.* [48], Pavlichin *et al.* [49] and Deorowicz *et al.* [50] to be only the first steps in this direction.

### Beyond storage and transfer

So far, we have discussed how compression can alleviate the burden of storing and transmitting various genomic data. It can, however, help in less obvious aspects as well.

One prominent example is *de novo* assembly for second-generation sequencing technology, based on the de Bruijn graph [55], where hundreds of GB of RAM could be needed with standard methods. Applying succinct data structures allowed to decrease the memory usage by an order of magnitude, and a whole assembly pipeline of a human individual was run in about 36 GB of RAM [56]. The nodes in the de Bruijn assembly graph are all distinct strings of some length  $k$  (e.g., 25) from given data, and edges between them exist if two nodes have a suffix-prefix overlap of length  $k - 1$ . The idea of Conway and Bromage [56] was to perceive the set of edges for given data as a subset of the full graph and to use (existing) succinct subset encoding techniques, supporting fast access. Representing this subset with naïve means would be impossible because of typically quadrillion-edge scale of the full graph. On the



other hand, standard approaches to the de Bruijn assembly graph construction are plagued with pointers (location addresses in the memory), which are dominant part of the data structure in RAM.

Compact data structures not always make use of “typical” compression techniques (like statistical coding or LZ-matches), yet they serve the same purpose, which is (in the currently discussed application) reducing the memory requirements for building or querying a large graph. The Bloom filters [57], the well-known idea for compact approximate subset representation, were recently used with success for the de Bruijn graph construction [58,59]. In particular, the work of Salikhov *et al.* [59] is a refinement of the technique of Chikhi and Rizk [58]. Their graph built for 564M human reads of length 100 bp using  $k = 23$  occupies only about 2.5 GB. Another succinct (but not exactly compression) technique was proposed by Ye *et al.* [60]. They store only a small subset of the observed  $k$ -mers as nodes, with their neighboring chains of bases as edges. Some care is taken to remove low-coverage edges, which normally result in tips, loops or bubbles in the assembly graph, undesirable in the graph for a few reasons, including very high memory requirements. The compact structure of SparseAssembler from the cited work was built in less than 2 hours for the human chromosome 14 with the peak memory use of 3 GB. The memory requirement for the whole human genome was below 30 GB. While the result from [59], cited above, seems clearly better, they are not directly comparable, unfortunately, due to different coverages and used values of  $k$ .

An alternative to the de Bruijn graph is the assembly *string graph* [61], not working on  $k$ -mers, but requiring fast and memory efficient algorithms for the computation of suffix-prefix overlaps of arbitrary length among reads. In a string graph, as opposed to the de Bruijn graph, each path represents a valid assembly of the reads (because the reads are not “decomposed” into many independent  $k$ -mers). Although this approach seems harder, in the SGA string assembler [62] error correction and assembling were performed with satisfactory accuracy on 125 Gbp of human genome reads using 54 GB of memory. This was achieved thanks to a compressed data structure, the FM-index [63], which will be mentioned also later. Interestingly, another practical string graph assembler, Readjoinder [64], which can process 115 Gbp short reads dataset in 52 GB of RAM, does not make use of compressed data structures, but its space effectiveness comes from an ingenious partitioning approach applied to the array of a relevant subset of all suffixes of all reads. Readjoinder also confirms that compact data structures may be fast because of locality of accesses to data.

Another compression example concerns data *indexing*. Computational biology is mostly about data analysis,

which in turn involves pattern search. If the data over which patterns are sought do not change over time, we talk about a static scenario. This is quite common, e.g., an already sequenced genome of a given individual is usually not updated for a long period. In such a case it may be worth to build an index structure for given data since its construction time, even if significant, is likely to be paid off during multiple subsequent pattern searches. A classic text indexing data structure is the *suffix tree* (ST) [43]. It is powerful and useful, but unfortunately requires up to  $28n$  bytes of space, where  $n$  is the sequence length. Thus, it is hard to store an ST in the main memory even for a single mammalian genome. The *compressed index* idea is to support all (or main) functionalities of its classic counterpart (e.g., returning the locations of all occurrences of the pattern in the text), but using much less space. The area of compressed indexes, initiated only around 2000, has been marked by tens of significant papers [65]. Unfortunately, those “general-purpose” text indexes are not a good choice for a collection of genomes of individuals of the same species. In these cases, LZ-based indexes are much more efficient in removing the specific (and very large) redundancy. Several works with LZ-style indexes designed for genomic data appeared in the recent years. Most of the solutions for the exact [66–68] and approximate pattern search [69] are rather theoretical as for only some of them implementations are available.

The Burrows–Wheeler transform (BWT) is used with huge success for mapping sequencing reads onto a reference genome, almost making the classic,  $q$ -gram-based approach, obsolete (for example, an interesting representative of the latter approach, Hobbes [70], is very fast but also uses large amount of memory). Some of the most important genome alignment algorithms, Bowtie [71,72], BWA [73], BWA-SW [74], SOAP2 [75], and GEM [76], make use of the FM-index [63] or another compressed index based on the BWT, occupying as little as about 2 GB for a human genome (with the exception of GEM, requiring usually from 3 GB to 6 GB). For more information on BWT and FM-index, see the Additional file 1. These aligners also belong to the fastest ones. All of them (Bowtie only in version 2) support ungapped and gapped alignments and all of them are multi-threaded to make use of multi-core CPUs. Using BWT for gapped alignment is cumbersome, and this is why Bowtie2 [72] and GEM combine it with dynamic programming, a classic computation technique boasting its flexibility and tolerance for large gaps and affine gap penalties. An important issue for compressed indexes is the working space needed during their construction, as standard BWT computation algorithms require at least  $5n$  bytes of memory. Lightweight algorithms for BWT computation [77,78] appeared relatively late, yet the Kärkkäinen’s method [77] is already implemented in Bowtie.

A special case of mapping sequence reads to genomes concerns RNA-Seq experiments, in which a 'snapshot' of RNA molecules in the cell is sequenced. The RNA-Seq [79] is a relatively new approach that proved highly successful especially for determination of gene expression. The main problem here is that we must deal with reads at exon-exon boundaries (without the introns present in the reference genome), so spliced mappings must be looked for, which is unusual for standard DNA reads mapping. Some mappers for this problem also make use of the FM-index, e.g., TopHat [80] and CRAC [81]. A comprehensive list of RNA-Seq mapping approaches can be found in [82].

The FM-index searches for a pattern finding its successive letters, from right to left, which can be called backward extension of a string. Recently, Li [83] presented a simple modification of the FM-index for forward-backward extension of DNA strings. His *de novo* assembler, fermi, shows that the assembly based variant calling can achieve an SNP accuracy close to the standard mapping approach, being particularly strong in indel calling.

It is worth noting that BWT-based alignment can be implemented on a massively parallel graphics processing unit (GPU). Recent tools SOAP3 [84] and SOAP3-dp [85], being about an order of magnitude faster than their CPU-based counterparts, are prominent examples.

One could ask if the FM-index, or another compressed index, is useful for searching DNA strings in a genome, e.g., given as FASTA input. The answer is positive; thanks to the small alphabet the search times (the *count* query, in which we return the number of matches only) of the FM-index, in the best current implementation, may be comparable with the suffix array ([86], Table Six and Seven). The space use, however, is only about  $0.3n$  (in contrast to  $5n$  needed by the suffix array). On the other hand, the *locate* query, in which the positions of all matching substrings are returned, is at least an order of magnitude slower, and needs some extra space.

In some cases, however, searching directly in the compressed data may be faster than in the straightforward representation. Loh *et al.* [87] compress a sequence database so that if an inserted sequence is similar enough to one from the database, it is represented as the reference plus a list of differences (edit script). The search algorithm they propose, based on BLAST, takes care of the differentially encoded sequences, and only rarely requires to bring them back to their "full" form. Their Compressive BLAST / BLAT algorithm was found to be about 4 times faster than classic BLAST / BLAT tools [88,89]. Other examples where data compression reportedly speeds up processing concern the *k*-mer counting task, especially in I/O-constrained scenarios ([90], Table Two–Four). We note that reading compressed input is nowadays a con-

venient feature of many tools (e.g., *de novo* assemblers Velvet [91] and ABySS [92]), but not always it brings improvements in speed.

Compression methods were used also for other purposes, in which the goal was not the reduction of space or processing speed-up, but rather better understanding of genomic data. Cao *et al.* [93] used the XM algorithm [37] to align eukaryotic-size genomes in a few hours on a workstation. The idea is to teach the expert models on one of the sequences and use the knowledge to properly align the second one by measuring the information content and the mutual information content of the sequences. The resulting aligner is shown experimentally to be superior (at least in quality, not in speed) to conventional alignment methods based on character matching.

Bhaduri *et al.* [94] proposed a somewhat related idea of using a compression algorithm from the LZ family to filter low-complexity reads in a project on identification of nonhuman sequences, such as viruses, in deep sequencing datasets.

A measure of sequence similarity, that is both accurate and rapidly computable, is highly desirable. Ferragina *et al.* [95] advocated that classic alignment methods do not scale well for huge data. They focus on the Universal Similarity Measure (USM) [96]. As USM is rather a theoretical concept, the authors experiment with its three approximations based on data compression. They validate the possibility of using these approximations for classification of sequences by UPGMA and NJ methods.

Freschi and Bogliolo [97] proposed a lossy compression scheme to eliminate tandem repeats from a sequence. Thanks to that, no repeat masking is necessary before performing pairwise alignment of sequences.

## Conclusions

Data deluge in computational biology has become a fact. A vast majority of gathered data is "temporary" in nature and could be discarded as soon as the analysis is done. The problem is, however, that current sequence analysis algorithms are imperfect, and storing lots of data only in the hope to squeeze out more of them in the future is a reasonable strategy. To put it in other words, lossy storage is an interesting option for bioinformatics, but it should be used judiciously.

The variety of genomic data formats implies the need for specialized compression algorithms better than the general-purpose standards, like gzip and bzip2. Succinct representation is not everything; decompression time or rapid access to arbitrary data snippets may matter even more, so they should be taken into account in algorithmic design. Sometimes even more enhanced functionalities are welcome. Fast search directly in the compressed data is an example. More efficient compression diminishes the costs of not only local data storage

and transfer, but also of data center services. The latter should bring the vision of ubiquitous cloud computing closer.

Let us make two predictions at the end. First, we note that some objects of interest in computational biology, like a human genome, do not grow. Hence, with growing amount of memory even in our home laptops, it perhaps no longer pays to apply strong compression for some tasks, if less compact but faster solutions are known. Read alignment onto a reference genome is a prominent example of this sort. We anticipate that in 1–2 years solutions processing a 1 billion 100 bp reads collection in a few hours on a PC will appear, but their main data structure may be the good old suffix array rather than, e.g., the FM-index.

Second, we predict that the turbulent period of new compression ideas for sequencing data representations will slowly give way to industry-oriented solutions, with more stress on robustness, flexibility, ease of use, and compression and decompression speed (in sequential and parallel/distributed regimes). Ideas are exiting, but routine jobs require standards. We believe that powerful, versatile and thus widely used formats in bioinformatics will emerge soon, proving the maturity of the field.

## Endnotes

<sup>a</sup> There are, of course, many alternative cloud storage solutions and it is hard to tell “typical” fees for storage and transfer, as opposed to retail disk media prices which can be monitored rather easily. As a reference, however, we note that Microsoft Windows Azure and Google Cloud Storage charges in the same scenario are similar (about 1,350–1,500 dollars), and all these providers charge more for the assumed 15 downloads than for one-year storage.

<sup>b</sup> Illumina software for their HiSeq 2500 equipment contains an option to reduce the number of quality scores [98]. Its effect on overall sequencing is shown in a technical support note [99].

<sup>c</sup> Statistical methods often encode symbols with regard to the gathered statistics of occurrences in their respective *contexts*, which are formed with, e.g., several proceeding symbols. This approach can be made even more sophisticated with considering several contextual models running in parallel, in order to improve the estimation of symbols’ probability and, in result, the obtained compression ratio. The name of “context mixing” refers to this approach, in which the statistics from different contexts are “mixed” (weighted, blended).

## Additional file

Additional file 1: Description of selected compression methods.

## Competing interests

The authors declare that they have no competing interests.

## Authors’ contributions

SD and SG contributed equally to the paper. Both authors wrote, read and approved the final manuscript.

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