A New Lossless DNA Compression Algorithm Based on A Single-Block Encoding Scheme

Deloula Mansouri, Xiaohui Yuan * and Abdeldjalil Saidani

School of Computer Science and Technology, Wuhan University of Technology, Wuhan 430070, China; deloula@whut.edu.cn (D.M.); saidani@whut.edu.cn (A.S.)
* Correspondence: yuanxiaohui@whut.edu.cn

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Abstract: With the emergent evolution in DNA sequencing technology, a massive amount of genomic data is produced every day, mainly DNA sequences, craving for more storage and bandwidth. Unfortunately, managing, analyzing and specifically storing these large amounts of data become a major scientific challenge for bioinformatics. Therefore, to overcome these challenges, compression has become necessary. In this paper, we describe a new reference-free DNA compressor abbreviated as DNAC-SBE. DNAC-SBE is a lossless hybrid compressor that consists of three phases. First, starting from the largest base (Bi), the positions of each Bi are replaced with ones and the positions of other bases that have smaller frequencies than Bi are replaced with zeros. Second, to encode the generated streams, we propose a new single-block encoding scheme (SEB) based on the exploitation of the position of neighboring bits within the block using two different techniques. Finally, the proposed algorithm dynamically assigns the shorter length code to each block. Results show that DNAC-SBE outperforms state-of-the-art compressors and proves its efficiency in terms of special conditions imposed on compressed data, storage space and data transfer rate regardless of the file format or the size of the data.

Keywords: DNA sequence; storage; lossless compression; FASTA files; FASTQ files; binary encoding scheme; single-block encoding scheme; compression ratio

1. Introduction

Bioinformatics is a combination of the field of informatics and biology, in which computational tools and approaches are applied to solve the problems that biologists face in different domains, such as agricultural, medical science and study of the living world.

DNA sequencing provides fundamental data in genomics, bioinformatics, biology and many other research areas. It contains some hidden significant genetic information, which makes it the most important element in bioinformatics and has become crucial for any biological or clinical research in different laboratories [1–3]. Genomic (or DNA) sequence is a long string formed of four nucleotide bases: adenine (A), cytosine (C), guanine (G) and thymine (T). This biomolecule not only contains the genetic information required for the functioning and development of all living beings but also aids in medical treatments in the detection of the risk for certain diseases, in the diagnosis of genetic disorders, and the identification of a specific individual [4,5]. Around three billion characters and more than 23 pairs of chromosomes are in the human genome, in a gram of soil there is 40 million bacterial cells, and certain amphibian species can even have more than 100 billion nucleotides [3,6]. With the emergent evolution in DNA sequencing technology, the volume of biological data has increased significantly.

Many file formats are used to store the genomic data such as FASTA, FASTQ, BAM/SAM and VCF/BCF [7–15]. In these file formats, in addition to the raw genomic or protein sequences, other information, such as in FASTQ file identifiers and quality scores are added. However, in FASTA
files only one identifier is added in the head of the data, which represents nucleotides or amino acids [16].

From 1982 to the present, the number of nucleotide bases generated in databases has doubled approximately every 18 months and their sizes can range from terabytes (TB) to petabytes (PB) [17]. In June 2019, approximately 329,835,282,370 bases were generated [18]. All of these data are stored in special databases, which have been developed by the scientific community such as the 1000 Genomes Project [6], the International Cancer Genome Project [19] and the ENCODE project [12,17,20]. This advancement has caused the production of a vast amount of redundant data at a much higher rate than the older technologies, which will increase in the future and may go beyond the limit of storage and bandwidth capacity [3–5]. Consequently, all these data pose many challenges for bioinformatics researchers, i.e., storing, sharing, fast-searching and performing operations on this large amount of genomic data become costly, requires enormous space, and has a large computation overhead for the encoding and decoding process [16]. Sometimes the cost of storage exceeds other costs, which means that storage is the primary requirement for unprocessed data [9].

To overcome these challenges in an efficient way, compression may be a perfect solution to the increased storage space issue of DNA sequences [3,21]. It is required to reduce the storage size and the processing costs, as well as aid in fast searching retrieval information, and increase the transmission speed over the internet with limited bandwidth [13,22]. In general, compression can be either lossless or lossy; in lossless compression, no information is lost and the original data can be recovered exactly, while in lossy compression, only an approximation of the original data can be recovered [23–26]. Many researchers believe that lossless compression schemes are particularly needed for biological and medical data, which cannot afford to lose any part of their data [3]. Thus, with looking at the importance of data compression, lossless compression methods are recommended for various DNA file formats such as FASTA and FASTQ file formats.

Currently, universal text compression algorithms, including gzip [27] and bzip [28], are not efficient in the compression of genomic data because these algorithms were designed for the compression of English text. Besides, the DNA sequences consist of four bases: two bits should be sufficient to store each base and follow no clear rules like those of text files that cannot provide proper compression results [9,29].

For example, bzip2 can compress 9.7 MB of data to 2.8 MB (the compression ratio (CR) is significantly higher than 2 bits per base (bpb)). Nevertheless, this is far from satisfactory in terms of compression efficiency. Thus, more effective compression techniques for genomic data are required.

Recently, several lossless compression algorithms have been suggested for the task at hand considering the features of these data.

Tahi and Grumbach introduced two general modes for DNA sequence lossless compression: horizontal mode and vertical mode [5]. The horizontal (or reference-free) methods are based only on the genetic information of the target sequences by referring exclusively to its substrings. Whereas the vertical (or reference-based) methods are using another genome sequence as a reference, and then store only the genomic locations and any mismatches instead of storing the sequences [29]. A significant advantage of referential compression is that as the read length increases, it produces much better compression ratios compared with non-referential algorithms. However, reference-based compression techniques are not self-contained and the success of these algorithms depends on the availability of a good reference genome sequence, which may not always be ensured. Additionally, the creation of such databases or reference sequences is a big challenging problem. In the decompression, the exact reference genome to match the genomic locations and extract the reads is required. Hence, it is important to preserve the reference genome [5].

Many studies have been proposed on the topic of DNA compression, considering the features of DNA sequences to achieve better performance, such as small alphabet size, repeats and palindromes. In general, these methods can be classified as statistical based, substitutional or dictionary based [13,29].
The first algorithm based substitutional compression found in the literature is Biocompress, which was developed in [30]. The replica of sequences, such as repeats, palindromes and complementary palindromes are detected using the Ziv and Lempel method, and then encoded them using the repeat length and the position of their earliest repeat occurrence. Two bits per base are used to encode non-repeat regions. BioCompress-2 [5] is an extension of BioCompress, exploiting the same methodology, as well as when no significant repetition is found, arithmetic coding of order-2 is applied. The use of “approximate repeats” in the compression of DNA sequences began with GenCompress [31], followed by its modified version DNACompress [32], DNAPack [16], GeNML [33] and DNAEcompress [34]. Two versions of GenCompress were proposed [31]. Hamming distance (only substitutions) is used to search approximate repeats in GenCompress-1 while the edition distance (deletion, insertion and substitution) is applied to encode the repeats in GenCompress-2. For non-repeat regions, arithmetic encoding of order 2 is used. However, GenCompress-2 fails to compress the larger DNA sequences. The DNACompress proposed in [32] consists of two phases. In the first phase, Pattern Hunter software [35] is used to identify the approximate repeats containing palindromes in the sequence. In the second phase, the Ziv and Lempel compression method is exploited to encode both approximate repeats and the non-repeat regions. In DNAPack [16], the authors found a better set of repeats than those found by the previous algorithms. By using a dynamic programming approach, the longest approximate repeats and the approximate complementary palindrome repeats are detected and then encoded by using the Hamming distance (only substitutions). Non-repeat regions are encoded by the best choice from context tree weighting, an order-2 arithmetic coding, and two bits per base. DNAPack performed better than the previous compressors in terms of compression ratio (with an average equal to 1.714 bpb using standard benchmark data).

The GeNML [33] presents the following improvements to the approach utilized in the NMLComp method [36]. First, it reduces the cost of the search for pointers of the previous matches by restricting the approximate repeats matches, then choosing the block sizes, which are utilized in parsing the target sequence. Finally, introducing adaptable overlooking elements for the memory model. Based on the statistical characteristics of the genomic sequence, the authors in [37] introduce the XM (eXpert Model) method that encodes each symbol by estimating its probability distribution based on the previous nucleotides utilizing a mixture of experts. Then, the results of experts are combined and encoded by arithmetic encoder. The DNASC algorithm [38] compresses the DNA sequence horizontally in the first step by using extended Lempel–Ziv style, and, then, vertically in the second step by taking different block size and window size.

In [4,39], two bits per base are used to encode the sequence of DNA as the bit-preprocessing stage. The GENBIT compress tool [39] divides the input DNA sequence into segments of eight bits (four bases). If the consecutive blocks are the same, a specific bit “1” is introduced as a 9th bit. Otherwise, the specific bit will be “0”. If the fragment length is less than four bases, then a unique two bits are assigned for each base. When the repetitive blocks are maximum, the best compression ratio for the entire genome is achieved. When the repetitive fragments are less, or nil, the compression ratio is higher than two. The DNABIT Compress Algorithm [4] splits the sequence into small fragments and compresses them while taking into consideration their existence before. To differentiate between exact and reverse repeat fragments, a binary bit in the preprocessing stage is used. This method achieves better compression and significantly improves the running time for larger genomes.

The concept of extended binary trees is used in the HuffBit Compress [40]. It replaced the bases A, C, G and T by 0, 10, 110 and 111, respectively. However, the compression ratio will be higher than two bits if the frequency of ‘G’ or ‘T’ is higher than other bases’ frequencies, which is the worst-case compression ratio results because both ‘G’ and ‘T’ are replaced by three bits. A combination of different text compression methods is proposed to compress the genomic data. The improved RLE, proposed in [41], is based on move-to-front (MTF), run length encoder (RLE) and delta encoding; this method is a good model to compress two or more DNA sequences.

The DNA-COMPACT (DNA COMpression) algorithm [42] based on a pattern-aware contextual modeling technique comprises of two phases. Firstly, it searched for exact repeats and palindromes by
exploiting complementary contextual models and represent them by a compact quadruplet. Secondly, where the predictions of these models are synthesized using a logistic regression model, then the non-sequential contextual models are used.

A new compression method named SBVRLDNAComp is proposed in [43]. In this method, the exact repeats are searched and then encoded in four different ways using different techniques, and then the optimal solution is applied to encode them. Finally, the obtained stream is compressed by LZ77. OBRLDNAComp [44] is a two-pass compression algorithm. In the first phase, it searches the optimal exact repeat within a sequence. Then, it scans a sequence horizontally from left to right followed by a vertical scanning from top to bottom before compressing them using a substitution technique. A seed-based lossless compression method for DNA sequence compaction, presented in [45], a new substitutional compression scheme similar to the Lempel-Ziv, is proposed to compress all the various types of repeats using an offline dictionary.

A combination of transformation methods and text compression methods was suggested in [46]. In the first step, Burrows–Wheeler transform (BWT) rearranges the data blocks lexicographically. MTF followed by RLE is used in the second step. Finally, arithmetic coding is used to compress the previously generated data. A novel attempt with the multiple dictionaries based on Lempel–Ziv–Welch (LZW) and binary search has been presented in [47]. The performance of this algorithm is tested using a non-real DNA dataset. To overcome the limitation of the Huffman algorithm [40]. A Modified Huffman compression algorithm was introduced in 2018 [48]. It encodes the bases according to their frequency, which means that the base that occurs more often will be encoded with the shortest code.

Recently, a cloud-based symbol table driven DNA compression algorithm (CBSTD) using R language was proposed in [6]. In this technique, the entire DNA sequence is subdivided into blocks (each block contains four bases), and then looked into which category it belongs. According to the occurrences of the nucleotide bases, three different categories are suggested. The first situation is when all four symbols are different, and hence, a table of 24 symbols is created to encode each block. The second situation is when the two nonconsecutive symbols are the same; then the block is divided into two parts and encoded using another table of symbols. The third situation is when two or more consecutive symbols are the same, in which case it is encoded with the bases followed by the number of occurrences. The authors in [6,40,48] do not use the real biological sequences to test the performance of their algorithms.

In [49], the authors illustrate an efficient genome compression method based on the optimization of the context weighting, abbreviated as OPGC. The authors in [50] introduced the OBComp, which is based on the frequencies of bases including other symbols than ACGT. Only a single bit is required to encode the two nucleotide bases that occur most frequently, and the positions of the two other bases will be recorded separately before removing them from the original file. The first output will be encoded using modified run length encoding [51] and Huffman coding [40], whereas the position files are transformed using data differencing, which can improve the final compression. One of the general compressors based on asymmetric binary coding is used to generate the final compressed data.

Generally, compression algorithms that look for approximate or even exact repeats have a high execution time overhead, as well as a high storage complexity, which is not suitable for an efficient compression technique. The following described methods are reference-free methods that have been designed for compressing files in FASTA and FASTQ formats.

Three of the selected tools compress FASTA files only. The first one is called BIND [52]. It is a lossless FASTA/multi FASTA compressor specialized for compressing nucleotide sequence data. After the separation of the head(s), 7-Zip is used to compress them; non-ATGC characters and lowercase character is also removed from the original data and their positions are stored separately. Each nucleotide (A, T, G and C) is represented with an individual binary code (00, 01, 10 and 11) and then splits the codes into two data streams. After that, the repeat patterns in these two data streams are encoded using a unary coding scheme. Finally, BIND uses the LZM algorithm to compress the output files.
The DELIMINATE [7] is an efficient lossless FASTA/multi FASTA compressor, it separately handles the header and sequence and it performs the compression in two phases. Firstly, all lower-case characters, as well as all non-ATGC, are recorded and eliminated from the original file. Secondly, the positions of the two most repeated nucleotide bases are transformed using delta encoding; then, the two other bases are encoded by binary encoding. Finally, all generated data output files are archived using 7-Zip.

The MFCompress [8] is one of the most efficient lossless non-referential compression algorithms for FASTA files compaction according to a recent survey [13]. It divides the data into two separate kinds of data: one containing the nucleotide sequences, the other one the headers of the FASTA records. The first data are encoded using probabilistic models (multiple competing finite context models) as well as arithmetic; whereas the headers are encoded using a single finite context models. The SeqCompress algorithm [9] is based on delta encoding with Lempel–Ziv; it uses a statistical model and arithmetic coding to compress DNA sequences.

The next generation sequencing data (NGS) are generally stored in FASTQ format. FASTQ files consist of millions to billions of records and each record has four lines, a metadata containing; a DNA sequence read obtained from one-fold of the over sampling; a quality score sequence estimating error probability of the corresponding DNA bases; and other line the same as the first one, which is generally ignored [1,5]. Due to the different nature of the FASTQ data, each compressor processed the various data (identifiers, DNA reads and quality score) separately. In this study, we are interested in the compression of DNA reads.

In [10], the authors present new FASTQ compression algorithm named DSRC (DNA sequence reads compressor). They impose a hierarchical structure of the compressed data by dividing the data into blocks and superblocks; it encodes the superblocks independently to provide fast random access to any record. Each block represents b records and each superblock contains k blocks (b = 32 and k = 512 by default). In this method, LZ77-style encoding, order-0 Huffman encoding is used for compressing the DNA reads. The Quip [53] is a reference-free compression scheme that use arithmetic coding based on order-3 and high order Markov chains for compressing all parts of FASTQ data. It also supports another file format such as the SAM and BAM.

In the Fqzcomp method [11], byte-wise arithmetic coder associated with order K context models are exploited for compressing the nucleotide sequence in the FASTQ file. As a reference free lossless FASTQ compressor, GTZ is proposed in [15]. It integrates adaptive context modeling with multiple prediction modeling schemes to estimate the base calls probabilities. Firstly, it transforms the DNA sequences with the BWT (Burrows–Wheeler transform) algorithm, which rearranges the sequences into runs of similar characters. Then, the appearance probability of each character is calculated by using the zero-order and the first-order prediction models. Finally, it compresses the generated data blocks with arithmetic coding.

In this paper, we propose a new lossless compression method DNAC-SEB that uses two different schemes to encode and compress the genomic data. Initially, based on the occurrence bases, the input data will be transformed into several binary streams using a binary encoding scheme. After that, all the binary streams generated previously are divided into fixed length blocks and encoded using the SEB method. This method encodes each block separately using two different techniques, and then the optimal code is chosen to encode the corresponding block. In the last step, one of a general-purpose compressor is used to generate the final compressed data. The proposed idea was inspired from [54,55].

The majority of these methods assume data drawn only from the ACGT alphabet, ignoring the appearance of other an unidentified base that can be found in DNA data sequences like N. However, the proposed approach considers that. The DNAC-SEB is a reference-free method; it does not depend on any specific reference genome or any patterns and may work with any type of DNA sequence and with no ATCG characters. It encodes each block of data separately and does not require any additional information in the decoding step.
The remaining part of this study is organized as follows. In the next section, we describe the compression method in detail. Then, we provide performance evaluation details and discussion of the proposed method against state-of-the-art DNA compressors using different benchmark. The last section presents the conclusion and some future work.

2. Proposed Methodology

2.1. Overview

The proposed algorithm works in three steps during compression and decompression, DNAC-SEB is a single-bit and single-block encoding algorithm based on the nucleotide base distribution in the input file. The general flowchart of the proposed method is shown in Figure 1.

Contrary to two-bit encoding methods, which assign two bits for each nucleotide base, the DNAC-SEB encodes each base using only one bit using binary encoding scheme, which depends on the nucleotide frequencies in the input file. Therefore, to improve the compression results, a new lossless compression method based on a single-block encoding scheme, named SBE, is proposed to encode all the previously generated binary stream. The main idea of this method is to split the binary stream into blocks with fixed length size (each block contains seven bits, because only two bits are used to represent the positions of zeros and ones within the blocks); it encodes each block separately without requiring any information or coding table by using the positions of 0’s and 1’s within the block. Then, the position of each bit within the block is calculated and two codes are generated based on the values of the bits zero or one. The shorter length code will be chosen to store the code of a respective block. All the processed data can be compressed with one of the existing compressors, such as Lpaq8.

The key components of the encoding and decoding process of the proposed schemes are discussed in the following section.

2.2. Binary Encoding Scheme

In this method, the DNA sequences were converted into several binary streams. Firstly, it counted the frequency (Fr) of each nucleotide in the input file and records it in descending order as B1, B2, B3, B4 and B5 occurrences (where B1 was assigned to the most frequent nucleotide and B5 was assigned to the base, which had the smallest count in the input data). Secondly, for N different nucleotide bases in the DNA sequences, N-1 binary files would be created to encode N-1 bases. Each of these files saved the positions of one specific base, i.e., File1 saved all the positions of B1, File2 saved all the positions of B2, etc., by replacing the positions of the base Bi with 1 and replacing the positions of others bases that have smaller frequencies than Bi with 0. The positions of bases with occurrences higher than Bi were ignored, the corresponding pseudocode was provided in Algorithm 1.
Algorithm 1. Binary Encoding Scheme

Begin
First phase (Binary Encoding Scheme)
Input: A DNA sequence \( \{b_1, b_2, b_3, \ldots, b_n\} \)
Output files: File_1, File_2, …, File_{n-1}.
Initialization: \( \text{File}_i(B_i) = \{} \)
Algorithm:
Let \( B_1, B_2, B_3 \) and \( B_4 \) is the descending frequencies order.
For each element of the sequence (j from 1 to n)
if \( (B_i = b_j) \)
Add 1 to \( \text{File}_i \)
else
if \( (B_i \neq b_j \text{ && } \text{Fr}(B_i) > \text{Fr}(b_j)) \)
Add 0 to \( \text{File}_i \)
else
Ignore it and pass to the next base
endif
endif
End.

It is important to note that the position of the base, which had the smallest occurring number, would be ignored because it could be recovered from the previously generated files.

2.3. Working Principle of the Single-Block Encoding Algorithm

The proposed algorithm is a single-block encoding that works on the principle of “positions based on zeros and ones”. It calculates the position of the current bit using the position of its previous neighbor bit(s) within block. The encoding scheme of the SBE is described in Figure 2.

Our algorithm for compressing the blocks worked as follows. First, we partitioned the binary input data into blocks with fixed length (in this case, each block contained seven bits). Initially, the position of each bit was calculated based on the value of the previous bit(s) and its position. After that, the SBE
started traversing from the second bit of the block to identify the bits with a zero value and save its position. Once a 0 bit was found, its position was stored in the code (code0) and the process was reinitiated until the last bit was attained. When the last bit was reached, the corresponding code would be stored. The first bit of the generated codes would identify which method was used, e.g., if 0-based position was used, then the first bit would be zero. After the completion of 0-based positions, the 1-based positions would take place and another code (code1) was obtained. Otherwise it would be one. The code of the lowest size was selected to encode the respective block. Finally, all the resultant optimal codes of encoded blocks were concatenated with the control bits, and the compressed file was generated. More details about that are explained in Figure 3.

![Flowchart](image)

**Figure 3.** The encoding block process using the single-block encoding scheme (SBE).

From Figure 3, the first bit of each block was recorded as a control bit and the procedure of the position was executed starting from the second bit, in which its position was always equal to zero. For each current bit (C), its position depends on the position(s) of its previous bit(s) (may be one bit or more). If C equals its previous adjacent bit (PA), then the position of C will be the same as the PA positions. Otherwise, if C is different from the PA, and the PA position is equal to one, then the position of C is also one. However, if C is different from the PA and the position of the PA is equal to 0, then we count the number of the previous bits (PPA) of the PA equal to the PA starting from the bit C until the PPA is different from the PA or the first bit is reached (the first bit is not included in the count), and then the counter value represents the position of C. All the obtained positions are recorded in position blocks that have the same size as the original blocks without counting the control bit. The proposed method is highly applicable to diverse DNA data.

In 0-based positions, once a value of zero is found, its position will be taken into consideration. Starting from the second bit, for each bit that equals zero and its position is equal to zero, then we add zero to the code. Further, if the position equals two, we add -00 to the code; if the position equals three,
then we add -01 to the code; and if the position equals four, we add -10 to the code. Finally, if the position equals five, then -11 is added to the code. When the last bit (seventh bit) is reached, the corresponding code will be stored as code-based 0s (code0). The process is reinitiated until the seventh block is reached.

After the completion of 0-based positions, the technique of 1-based positions will take place to generate the second code for the same block (code1). The process used in 0-based positions and 1-based positions is the same except that in the first process we search the positions of zeros in the binary digits; whereas in the second one we will find the positions of ones in the same block. Once code0 and code1 are generated, the proposed algorithm compares them and chooses the shorter one that had the minimum number of bits as optimal code (a special symbol used to encode the position). The number of bits in optimal code may contain a minimum of two bits and a maximum of six bits. Finally, all the resultant optimal codes of encoded blocks are concatenated to generate the compressed file.

2.4. Illustration

To further understand the proposed method, an illustration is given here. Let us consider an input sequence (D) with (D) = {TACTGNTAAAGTACNATTGNTCTAGNCTACCCGCA}, of length k (k = 40), which contains A, C, T, G and N bases. The proposed approach operates as follows. In the first step, the frequency of the existed bases was counted and the bases were recorded in descending order. So, ATCGN was the descending order of the bases, in which 13, 9, 8, 6 and 4 were the frequencies of A, T, C, G and N, respectively. Then, four files were generated in this step, which is more detailed in Figure 4.

![Figure 4. The encoding process of the binary encoding scheme.](image-url)

From the Figure, File1, File2, File3 and File4 were generated using the first proposed method, which presented the binary encoded positions of A, T, C and G, respectively. The positions of the unidentified base N were ignored because it could be recovered from the output files. Each of these files was divided into blocks of seven bits and each block was processed as follow. The workflow of the first block of File1 is illustrated in the Figure 5 above.

Let us consider the previously generated binary streams as an input sequence of the SEB scheme. Each binary sequence was divided into a block of seven bits, so the encoding process of the first block of this sequence using the SBE scheme was as follows.

The first bit in the block was 0, so 0 was recorded as the control bit. The position of each bit in the block was calculated (see Figure 4), the position of the second bit (0100000) was considered as 0th for all situations, and the positions block was written as 0 - - - - - . The third bit of the sequence (0100000) would be compared with its previous bits; 0 was different to 1 and the second bit was processed; then the position of this bit would be 1 and the positions block was written as 01 - - - - . For the fourth bit (0100000), its value was equal to its previous neighbor value (the third bit); so, its position would be 0 and the position block was written as 010 - - - . In the same way, the position of the fifth bit was calculated, the position block was written as 0100 - - . If the value of the sixth bit was the same as its previous one, then its position would be 0 and the position block was written as 01000 - . The position of the last bit was also equal to its first previous bit, so the obtained position block was 010000.

Using 0-based positions, the first bit of code0 would be 0 adding the positions of 0s in the obtained position block, the generated code was 01000. The same method was used for 1-based positions, but just the position of 1 would be taken into consideration, so the generated code1 was 10. The shorter code
in terms of number of bits was selected to encode the block, so 0100000 would be encoded with 010. Finally, the encoded the first block 0100000 was successfully decoded in the decompression process. This process repeated for every block in the input sequence and the overall input file was reconstructed with no loss of information.

![Workflow of the encoding process of the first block using the SEB scheme.](image)

**Figure 5.** Workflow of the encoding process of the first block using the SEB scheme.

### 2.5. Decoding Process

The proposed approach was based on lossless compression, which means that the decompression process was exactly the reverse operation as applied in the compression process. No additional information is needed to be transmitted along with the compressed data. The input data before compression and the output data after decompression were exactly the same. Thus, this phase was exactly the reverse operation of the encoding phase and also consisted of three steps; first, we applied the lpaq8 decompressor; secondly, each bit in the control bits file represents the first bit of each block and each first bit of each code indicate which method was used (positions based on 1s or based on 0s). When it is based on 0s, the positions of 0s in the code were replaced by 0 and the remaining places were filled with 1, removing the first bit of this block. In the same way, if the code was based on 1s, the positions in this code were represented by 1 and the remaining positions were filled with 0 and also the first bit of the code would be removed. The block was reconstructed from the concatenation of the corresponding control bit and the generated code (with six bits). After that, for each generated file binary decoding algorithm was applied based on the descending order of the nucleotide bases began the compression from the highest base count. Then, the original sequence was constructed with no loss of information. Let us consider the first block, the shorter code block was 010, the decoding process of the first compressed block is illustrated in Figure 6.

The first bit represents the control bit; then the decoded sequence would be 0 - - - - - - . The first bit of the optimal code was 1 (10), which means that the positions in the code would be replaced with 1. The second bit in the code (10) indicates the position 0 with respect to the first bit, and it should be populated with 1. It implies that the value of the second bit should be populated with 1, and the decoded sequence would be 01 - - - - - . This code contained only one position, which indicates that the 2nd, 3rd, 4th and 6th bits should be represented by zeros, which would generate the original block 0100000.
We also carried out tests on seven publicly accessible FASTA sequences downloaded from sequence read archive (SRA) \cite{56}. Tables 1–3 provide the descriptions of the datasets.

The data sets include the KOREF 2009024, NC_017526, NC_017652, sacCer3, chimpanzee, \textit{Caenorhabditis elegans}, \textit{Clostridium symbiosum} and others ranging from 38 to 223 KB taken from a variety of sources with different species, types and sizes were chosen. The first one included diverse DNA datasets (chloroplast DNA, human DNA, virus DNA, mitochondria DNA, \textit{Pseudomonas}, \textit{Caenorhabditis elegans}, \textit{Clostridium symbiosum} and others) ranging from 38 to 223 KB taken from a variety of sources with different, which were downloaded from the National Centre for Biotechnology Information (NCBI) database \cite{56}. We also carried out tests on seven publicly accessible FASTA/multi FASTA data sets, which have already been used in various studies and available in different databases such as NCBI and UCSC website. The data sets include the KOREF 2009024, NC_017526, NC_017652, sacCer3, chimpanzee, eukaryotic and C.e10 dataset. These datasets consist of FASTA files varying in size from a few kilobytes to megabytes. The third dataset contained four real-world DNA reads extracted from FASTQ data obtained from various high-throughput DNA sequencing platforms ranging from 28.7 to 1536 MB were downloaded from sequence read archive (SRA) \cite{56}. Tables 1–3 provide the descriptions of the above datasets.

3. Performance Evaluation

We benchmarked the proposed method against state-of-the-art reference-free genomic data compressors. The dataset proposed for this benchmark contained three different DNA file formats, with a consistent balance between the number of files, sequences and sizes. The dataset contained a total DNA sequence length of 9,420,117,815 bases (9 gigabytes).

For making the comparison, all of the compression methods were executed under a Linux OS; then, their performance was evaluated in terms of C-Size (MB), C-ratio (bpb), C-Mem, D-Mem (MB), C-Time and D-Time (s) stand for the compressed size, compression ratio, compression and decompression memory usage in megabytes, compression and decompression times in seconds, respectively. The results of compressing the several genomic data sequences using different compression metrics are shown with a bold font. It is important to note that our algorithm accepted data from all types of platforms and designed only to compress DNA sequences and did not accept any other input data.

3.1. Dataset Used

To ensure the comprehensiveness of our evaluation, three file formats of DNA datasets with different species, types and sizes were chosen. The first one included diverse DNA datasets (chloroplast DNA, human DNA, virus DNA, mitochondria DNA, \textit{Pseudomonas}, \textit{Caenorhabditis elegans}, \textit{Clostridium symbiosum} and others) ranging from 38 to 223 KB taken from a variety of sources with different, which were downloaded from the National Centre for Biotechnology Information (NCBI) database \cite{56}. We also carried out tests on seven publicly accessible FASTA/multi FASTA data sets, which have already been used in various studies and available in different databases such as NCBI and UCSC website. The data sets include the KOREF 2009024, NC_017526, NC_017652, sacCer3, chimpanzee, eukaryotic and C.e10 dataset. These datasets consist of FASTA files varying in size from a few kilobytes to megabytes. The third dataset contained four real-world DNA reads extracted from FASTQ data obtained from various high-throughput DNA sequencing platforms ranging from 28.7 to 1536 MB were downloaded from sequence read archive (SRA) \cite{56}. Tables 1–3 provide the descriptions of the above datasets.

![Figure 6.](image) The decoding process of the first compressed code block.
Table 1. A selected DNA dataset.

| Sequence   | Description                                                                 | Source            | Uncompressed Size (bytes) |
|------------|------------------------------------------------------------------------------|-------------------|--------------------------|
| CHMPXX     | Marchantia polymorpha chloroplast genome DNA                                | Chloroplasts      | 121,024                  |
| CHNTXX     | Nicotiana tabacum chloroplast genome DNA                                     |                   | 155,844                  |
| HUMHBB     | Human beta globin region on chromosome 11.                                  |                   | 73,308                   |
| HUMGHCSA   | Human growth hormone (GH1 and GH2) and chlorionic somatomammotropin (CS-1, CS-2 and CS-5) genes, complete cds. | Human             | 66,495                   |
| HUMHPRTB   | Human hypoxanthine phosphoribosyl transferase gene, complete cds.           |                   | 56,737                   |
| HUMDYSTROP | Homo sapiens dystrophin (DHD) gene, intron 44                               |                   | 38,770                   |
| MTPACGA    | Marchantia polymorpha mitochondrion, complete genome.                       | Viruses           | 100,314                  |
| MPOMTCG    | Marchantia polymorpha mitochondrion, complete genome.                       |                   | 186,609                  |
| HEHCMVCG   | Human cytomegalovirus strain AD169 complete genome.                         | Mitochondria      | 229,354                  |
| VACC        | Vaccinia virus, complete genome.                                            |                   | 191,737                  |

Table 2. A selected FASTA genome dataset.

| Genome     | Species                 | Size (bytes) |
|------------|-------------------------|--------------|
| KOREF 2009024 | Homo sapiens            | 3,131,781,939|
| NC 017652   | E. coli                 | 5,106,565.12  |
| NC_017526   | L. pneumophila          | 2,715,811.84  |
| Ce10        | C. elegans              | 102,288,589   |
| sacCer3     | S. cerevisiae           | 12,384,168.3  |
| Eukaryotic  | Haploid set of human chromosomes | 2,509,353,796 |
| Chimpanzee  | Pan troglodytes         | 3,333,423,104 |

Table 3. A selected DNA-FASTQ dataset.

| Data       | Species/Type            | DNA Sequence Size (bytes) |
|------------|-------------------------|---------------------------|
| ERR005143_1 | P. syringae            | 127,840,788               |
| ERR005143_2 |                         | 127,840,788               |
| SRR554369_1 | Pseudomonas aeruginosa | 165,787,101               |
| SRR554369_2 |                         | 165,787,101               |

3.2. Compression Performance

The performance of our algorithm was evaluated in terms of compression ratio (CR), memory usage, as well as compression and decompression time. CR was obtained by dividing the number of bits in compressed data by the number of bits in the original data [52].

To assess the efficiency of the proposed method, its compression performance was compared with the other lossless DNA compressors, namely Biocompress-2 [5], DNACompress [32], DNAPack [16], improved RLE [41], DNA-COMPACT (DNA-COMP) [42], Modified Huffman [48] and others in terms of CR using a real DNA dataset. Moreover, FS4CP [57] and OPGC [49], as well as the OBComp algorithm [50] were also included in the comparison. Traditional compression algorithms such as gzip [27], bzip2 [28] and arithmetic [47] coding and some of the transformation techniques such as BWT and MTF were also used to compress the DNA sequences. The obtained compression ratios of the DNAC-SEB against state-of-the-art algorithms are shown in Tables 3 and 4. It should be noted that in Tables 4–9, the best results in terms of different compression metrics are shown with a bold font.

Based on Table 4, the proposed technique had a compression ratio as good as the DNA compression method. Note that data pertaining to HUMGHCS, CHMPXX and MTPACGA show excellent results, in which the size was reduced by more than 78%, in particular using DNAC-SBE.
Regarding arithmetic coding and the results shown on the Table 5, the method provided a small factor of compressibility, better than bzip2 5.92%. At the same time, context tree weighting (CWT) [58] managed to perform better than other methods, except our algorithm. The DNAC-SBE outperformed other text compression algorithms on all the applied DNA datasets. The combination of different transformation methods also failed to compress the DNA sequence efficiently because the DNA sequences followed no clear rules like those of text data, which could not provide proper compression results.

Table 5. Performance comparison between various text compression algorithms and some of Burrows–Wheeler transform (BWT) based algorithms in terms of compression ratio (CR).

| Sequence | HEHC-MVCG | HUM-HBB | VAC-CG |
|----------|-----------|---------|--------|
| gzip     | 1.85      | 2.23    | 2.19   |
| Bzip2    | 2.17      | 2.15    | 2.10   |
| Arithm.  | 2.05      | 2.12    | 1.86   |
| CTW      | 1.84      | 1.92    | 1.86   |
| BWT+MTF+RLE+arithm. | 2.24 | 2.21 | 2.18 |
| BWT+MTF+arithm. | 2.03 | 2.02 | 2.07 |
| BWT+MTF  | 2.02      | 2.00    | 1.95   |
| DNAC-SBE | 1.74      | 1.72    | 1.67   |

The percentage reduction of compressing genomic FASTA data sequences using DNAC-SEB and different methods (that do not reorder the input sequences) in comparison to 7z are reported in Table 6. Original file sizes and compressed data sizes in megabytes with 7z are also shown in columns 2 and 3 in the same table.

From the results, we observed that the percentage reduction attained by DNAC-SEB was higher than Bind, Deliminate, MFCompress and SeqCompress. DNAC-SEB gave the best results for all the used datasets and SeqCompress gave the second to best percentage reduction for all datasets except for the eukaryotic. Other aspects result such as compressed size, execution time and memory usage using different FASTA genomic data sequences are reported in Table 7.
Table 6. Percentage reduction attained by Bind, Deliminate, MFCompress and SeqCompress in comparison to 7z. The sizes of compressed data using 7Z are shown in (MB).

| Dataset       | Original Size | 7z       | Bind   | Deliminate | MFCompress | SeqCompress | DNAC-SEB |
|---------------|---------------|----------|--------|------------|------------|-------------|----------|
| NC_017526     | 2.59          | 0.72     | 11.11% | 12.5%      | 9.72%      | 16.67%      | 23.61%   |
| NC_017652     | 4.87          | 1.36     | 11.76% | 12.5%      | 10.29%     | 16.91%      | 22.06%   |
| Ce10          | 97.55         | 27.81    | 5.11%  | 14.42%     | 17.12%     | 18.62%      | 28.98%   |
| sacCer3       | 11.82         | 3.33     | 16.68% | 19.6%      | 22.08%     | 23.44%      | 25.53%   |
| Eukaryotic    | 2139.98       | 527.27   | 11.31% | 15.44%     | 17.35%     | 17.11%      | 19.71%   |

Table 7. Comparison of compressed data size, processing time and memory utilization using FASTA datasets.

| Dataset       | Method     | Metrics    | C-Size | C-Mem | D-Mem | C-Time | D-Time |
|---------------|------------|------------|--------|-------|-------|--------|--------|
| NC_017526     | Bind       |            | 0.64   | 3.67  | 8.6   | 0.64   | 0.26   |
|               | Deliminate |            | 0.63   | 3.66  | 6.55  | 0.37   | 0.23   |
|               | MFCompress |            | 0.65   | 514.16| 513.13| 0.66   | 0.55   |
|               | DNAC-SEB   |            | 0.55   | 36.71 | 64.31 | 1.78   | 1.24   |
| NC_017652     | Bind       |            | 1.2    | 4.04  | 8.6   | 0.49   | 0.36   |
|               | Deliminate |            | 1.19   | 4.02  | 6.55  | 0.44   | 0.25   |
|               | MFCompress |            | 1.22   | 514.34| 513.13| 1.26   | 0.95   |
|               | DNAC-SEB   |            | 1.06   | 49.03 | 101.45| 1.67   | 1.27   |
| sacCer3       | Bind       |            | 3.16   | 5.75  | 8.12  | 2.78   | 2.41   |
|               | Deliminate |            | 2.85   | 2.56  | 6.19  | 5.26   | 4.94   |
|               | MFCompress |            | 2.77   | 473.9 | 473.78| 4.32   | 3.5    |
|               | DNAC-SEB   |            | 2.48   | 5.09  | 13.54 | 26.57  | 9.71   |
| Ce10          | Bind       |            | 23.17  | 37.92 | 7.54  | 11     | 7.27   |
|               | Deliminate |            | 22.36  | 70.29 | 7.11  | 8.55   | 8.54   |
|               | MFCompress |            | 21.67  | 433.92| 431.49| 20.16  | 19.04  |
|               | DNAC-SEB   |            | 19.75  | 68.75 | 212.75| 73.13  | 45.04  |
| Chimpanzee    | Bind       |            | 619.35 | 274.48| 28.52 | 350.84 | 180.28 |
|               | Deliminate |            | 590.48 | 284.02| 29.44 | 257.02 | 172.83 |
|               | MFCompress |            | 581.12 | 482.8 | 465.12| 575.22 | 426.53 |
|               | DNAC-SEB   |            | 570.73 | 522.23| 792.9 | 1445.15| 1155.97|
| Korea2009024  | Bind       |            | 693.14 | 295.05| 30.06 | 380.38 | 207.44 |
|               | Deliminate |            | 604.16 | 305.41| 31.07 | 303.15 | 194.99 |
|               | MFCompress |            | 604.12 | 665.44| 577.45| 503.18 | 500.31 |
|               | DNAC-SEB   |            | 577.23 | 649.56| 703.69| 1479.57| 1003.27|
| Eukaryotic    | Bind       |            | 466.98 | 387.81| 39.12 | 285.24 | 154.01 |
|               | Deliminate |            | 445.89 | 389.1 | 40.3  | 206.43 | 136.83 |
|               | MFCompress |            | 435.79 | 478.86| 477.89| 468.18 | 419.15 |
|               | DNAC-SEB   |            | 423.32 | 1859  | 1349.05| 1117.44| 903.1  |

From the result shown in Table 7, in all cases, the proposed approach generated a small file size as compared to other methods and hence provided the best results in terms of compression ratio. This was due to the modeling step and SEB method, which made the input files more suitable for the final compression. However, generally Deliminate uses less memory in both compression and decompression rather than other compressors. Since Bind was the fastest one in compression and decompression, it yielded a worse compression ratio in comparison to other competitors. To assess the efficiency of the proposed approach, its compression performance was also compared to some of FASTQ compression methods in terms of CR, execution time and memory usage. The obtained results are tabulated in Tables 8 and 9.
From Table 8 values, it is clear that the proposed method achieved a better compression ratio than other methods on all the applied dataset. In all cases, DNAC-SEB generated low file size as compared to other algorithms and hence a better compression ratio. Using ERR005143_1 dataset, DNAC-SEB shows an improvement of 4.78% and 3.52% to GTZ and Fqzcomp, respectively. The FASTQ file compressors, including Fqzcomp [11], Quip [53], DSRC [10] and gzip [27], were compared and the results are reported in Table 9.

Table 9. Comparison of compressed data size, processing time and memory utilization using FASTQ datasets.

| Dataset       | Method    | C-Size | C-Mem | D-Mem | C-Time | D-Time |
|---------------|-----------|--------|-------|-------|--------|--------|
| ERR005143_1   | DSRC      | 30.46  | 166.45| 100.56| 5.54   | 3.4    |
|               | Quip      | 28.96  | 393.7 | 389.02| 10.63  | 22.09  |
|               | Fqzcomp   | 27.73  | 79.46 | 67.33 | 7.88   | 10.08  |
|               | Gzip      | 124.36 | 0.86  | 1.39  | 28.94  | 2.7    |
|               | DNAC-SEB  | 26.81  | 580.38| 563.91| 38.08  | 18.64  |
| ERR005143_2   | DSRC      | 30.41  | 168.56| 106.5 | 5.83   | 4.09   |
|               | Quip      | 29.25  | 395.25| 389.54| 11.07  | 23.95  |
|               | Fqzcomp   | 27.9   | 79.46 | 59.3  | 8.45   | 10.35  |
|               | Gzip      | 132.42 | 0.86  | 1.39  | 29.45  | 2.79   |
|               | DNAC-SEB  | 26.28  | 580.57| 524.57| 72.06  | 20.54  |
| SRR554369_1   | DSRC      | 39.52  | 183.42| 152.65| 4.11   | 4.46   |
|               | Quip      | 35.26  | 383.95| 382.59| 10.86  | 22.93  |
|               | Fqzcomp   | 35.13  | 79.47 | 67.34 | 7.21   | 10.4   |
|               | Gzip      | 45.97  | 0.87  | 1.39  | 28.51  | 1.05   |
|               | DNAC-SEB  | 34.9   | 702.18| 615.97| 43.71  | 31.34  |
| SRR554369_2   | DSRC      | 39.38  | 173.57| 132.1 | 4.23   | 4.6    |
|               | Quip      | 35.93  | 383.84| 382.1 | 11.63  | 23.62  |
|               | Fqzcomp   | 35.12  | 66.41 | 57.179| 7.86   | 10.52  |
|               | Gzip      | 45.89  | 0.87  | 1.39  | 27.98  | 1.05   |
|               | DNAC-SEB  | 34.84  | 781.98| 644.66| 43.18  | 30.8   |

The results of FASTQ file compression with different compressors show that DNAC-SEB outperformed existing tools in terms of compression ratios for all types of FASTQ benchmark datasets. From the experimental results, we could observe that the best compression ratio, minimum memory and maximum speed requirements were competing goals. Generally, the higher compression rates were achieved by compressors that were slower and required higher memory. For example, Gzip had the best performance in the case of memory and decompression speed usage but offered the lowest compression ratio, which provided valuable long-term space and bandwidth savings. When the data size was an important aspect, tools such as DNAC-SEB were crucial.

4. Conclusions

In this paper, the authors described a new reference-free DNA compressor (DNAC-SEB), a lossless hybrid compressor that consisted of three phases, to compress any type of DNA sequences. The obtained
results show that DNAC-SBE outperformed some of the state-of-the-art compressors on the benchmark performed in terms of the compression ratio, regardless of the file format or data size. The proposed algorithm compressed both repeated and non-repeated sequences, not depending on any specific patterns or parameters and accepting data from all kinds of platforms. An important achievement of the paper was that it took into consideration the appearance of an unidentified base found in DNA data sequences, ignored by most of the other compression methods. Another advantage of this approach was that the DNAC-SEB compressor guaranteed the full recovery of the original data. The design of DNAC-SBE was simple and flexible, showing good compression rates compared to an important part of other similar algorithms.

DNAC-SEB is more suitable when the compressed data, storage space and data transfer were crucial. More efforts will be made in the future to optimize memory utilization and compression speed.

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**Availability of Data and Materials:** Some of the datasets cited used in this study were downloaded from NCBI sequence read archive (SRA) from the URL ftp://ftp.ncbi.nih.gov/genomes/. And some of them are obtained from UCSC database from the URL: ftp://hgdownload.soe.ucsc.edu/goldenPath. The FASTQ datasets were download from the URL: ftp://ftp.sra.ebi.ac.uk/vol1/fastq/.

**References**

1. Lander, E.S.; Linton, L.M.; Birren, B.; Nusbaum, C.; Zody, M.C.; Baldwin, J.; Devon, K.; Dewar, K.; Doyle, M.; FitzHugh, W.; et al. Initial sequencing and analysis of the human genome. *Nature* **2001**, *409*, 860–921. [PubMed]
2. Saada, B.; Zhang, J. Vertical DNA sequences compression algorithm based on hexadecimal representation. In Proceedings of the World Congress on Engineering and Computer Science, San Francisco, CA, USA, 21–23 October 2015; pp. 21–25.
3. Jahaan, A.; Ravi, T.; Arokiaraj, S. A comparative study and survey on existing DNA compression techniques. *Int. J. Adv. Res. Comput. Sci.* **2017**, *8*, 732–735.
4. Rajarajeswari, P.; Apparao, A. DNABIT Compress–Genome compression algorithm. *Bioinformation* **2011**, *5*, 350. [CrossRef] [PubMed]
5. Grumbach, S.; Tahi, F. A new challenge for compression algorithms: Genetic sequences. *Information Process. Manag.* **1994**, *30*, 875–886. [CrossRef]
6. Majumder, A.B.; Gupta, S. CBSTD: A Cloud Based Symbol Table Driven DNA Compressions Algorithm. In *Industry Interactive Innovations in Science, Engineering and Technology*; Springer: Singapore, 2018; pp. 467–476.
7. Mohammed, M.H.; Dutta, A.; Bose, T.; Chadaram, S.; Mande, S.S. DELIMINATE—a fast and efficient method for loss-less compression of genomic sequences: Sequence analysis. *Bioinformatics* **2012**, *28*, 2527–2529. [CrossRef]
8. Pinho, A.J.; Pratas, D. MFCompress: A compression tool for FASTA and multi-FASTA data. *Bioinformatics* **2014**, *30*, 117–118. [CrossRef] [PubMed]
9. Sardaraz, M.; Tahir, M.; Ikrar, A.A.; Bajwa, H. SeqCompress: An algorithm for biological sequence compression. *Genomics* **2014**, *104*, 225–228. [CrossRef]
10. Deorowicz, S.; Grabowski, S. Compression of DNA sequence reads in FASTQ format. *Bioinformatics* **2011**, *27*, 860–862. [CrossRef]
11. Bonfield, J.K.; Mahoney, M.V. Compression of FASTQ and SAM format sequencing data. *PloS ONE* **2013**, *8*, e59190. [CrossRef]
12. Aly, W.; Yousuf, B.; Zohdy, B. A Deoxyribonucleic acid compression algorithm using auto-regression and swarm intelligence. *J. Comput. Sci.* 2013, 9, 690–698. [CrossRef]

13. Hosseini, M.; Pratas, D.; Pinho, A.J. A survey on data compression methods for biological sequences. *Information* 2016, 7, 56. [CrossRef]

14. Numanagić, I.; Bonfield, J.K.; Hach, F.; Voges, J.; Ostermann, J.; Alberti, C.; Mattavelli, M.; Sahinalp, S.C. Comparison of high-throughput sequencing data compression tools. *Nat. Methods* 2016, 13, 1005. [CrossRef]

15. Xing, Y.; Li, G.; Wang, Z.; Feng, B.; Song, Z.; Wu, C. GTZ: A fast compression and cloud transmission tool optimized for FASTQ files. *BMC bioinformatics* 2017, 18, 549. [CrossRef] [PubMed]

16. Behzadi, B.; Le Fessant, F. DNA compression challenge revisited: A dynamic programming approach. In Proceedings of the Annual Symposium on Combinatorial Pattern Matching, Heidelberg, Jeju Island, Korea, 19–22 June 2005; pp. 190–200.

17. Kuruppu, S.; Puglisi, S.J.; Zobel, J. Reference sequence construction for relative compression of genomes. In Proceedings of the International Symposium on String Processing and Information Retrieval, Pisa, Italy; 2011; pp. 420–425.

18. GenBank and WGS Statistics (NCBI). Available online: https://www.ncbi.nlm.nih.gov/genbank/statistics/ (accessed on 29 June 2019).

19. 1000 Genomes Project Consortium “a map of human genome variation from population-scale sequencing”, *Nature* 467 (2010) 1061–1073. Available online: www.1000genomes.org (accessed on 19 April 2020).

20. Consortium, E.P. The ENCODE (ENCyclopedia of DNA elements) project. *Science* 2004, 306, 636–640. [CrossRef] [PubMed]

21. Keerthy, A.; Appadurai, A. An empirical study of DNA compression using dictionary methods and pattern matching in compressed sequences. *IJAR* 2015, 10, 35064–35067.

22. Arya, G.P.; Bharti, R.; Prasad, D.; Rana, S.S. An Improvement over direct coding technique to compress repeated & non-repeated nucleotide data. In Proceedings of the 2016 International Conference on Computing, Communication and Automation (ICCCA), Noida, India, 29–30 April 2016; pp. 193–196.

23. Rastogi, K.; Segar, K. Analysis and performance comparison of lossless compression techniques for text data. *Int. J. Eng. Comput. Res.* 2014, 3, 123–127.

24. Singh, A.V.; Singh, G. A survey on different text data compression techniques. *Int. J. Sci. Res.* 2012, 3, 1999–2002.

25. Al-Okaily, A.; Almarri, B.; Al Yami, S.; Huang, C.-H. Toward a Better Compression for DNA Sequences Using Huffman Encoding. *J. Comput. Biol.* 2017, 24, 280–288. [CrossRef]

26. Sharma, K.; Gupta, K. Lossless data compression techniques and their performance. In Proceedings of the 2017 International Conference on Computing, Communication and Automation (ICCCA), Greater Noida, India, 5–6 May 2017; pp. 256–261.

27. Gzip. Available online: http://www.gzip.org/ (accessed on 29 June 2019).

28. Bzip. Available online: http://www.bzip.org/ (accessed on 29 June 2019).

29. Bakr, N.S.; Sharawi, A.A. DNA lossless compression algorithms. *Am. J. Bioinformatics Res.* 2013, 3, 72–81.

30. Grumbach, S.; Tahi, F. Compression of DNA sequences. In Proceedings of the Data Compression Conference (DCC-93), Snowbird, UT, USA, 30 March–2 April 1993; pp. 340–350.

31. Chen, X.; Kwong, S.; Li, M. A compression algorithm for DNA sequences and its applications in genome comparison. *Genome Inf.* 1999, 10, 51–61.

32. Chen, X.; Li, M.; Ma, B.; Tromp, J. DNACompress: Fast and effective DNA sequence compression. *Bioinformatics* 2002, 18, 1696–1698. [CrossRef] [PubMed]

33. Korodi, G.; Tabus, I. An Efficient Normalized Maximum Likelihood Algorithm for DNA Sequence Compression. *ACM Trans. Inf. Syst.* 2005, 23, 3–34. [CrossRef]

34. Tan, L.; Sun, J.; Xiong, W. A Compression Algorithm for DNA Sequence Using Extended Operations. *J. Comput. Inf. Syst.* 2012, 8, 7685–7691.

35. Ma, B.; Tromp, J.; Li, M. PatternHunter—Faster and more sensitive homology search. *Bioinformatics* 2002, 18, 440–445. [CrossRef]

36. Tabus, I.; Korodi, G.; Rissanen, J. DNA sequence compression using the normalized maximum likelihood model for discrete regression. In Proceedings of the Data Compression Conference (DCC ’03), Snowbird, UT, USA, 25–27 March 2003; pp. 253–262.
37. Cao, M.D.; Dix, T.I.; Allison, L.; Mears, C. A simple statistical algorithm for biological sequence compression. In Proceedings of the 2007 Data Compression Conference (DCC'07), Snowbird, UT, USA, 27–29 March 2007; pp. 43–52.

38. Mishra, K.N.; Aggarwal, A.; Abdelhadi, E.; Srivastava, D. An efficient horizontal and vertical method for online DNA sequence compression. Int. J. Comput. Appl. 2010, 3, 39–46. [CrossRef]

39. Rajeswari, P.R.; Apparao, A.; Kumar, R.K. Huffbit compress—Algorithm to compress DNA sequences using extended binary trees. J. Theor. Appl. Inform. Tech. 2010, 10, 101–106.

40. Rajeswari, P.R.; Apparao, A.; Kumar, R.K. Huﬀbit compress—Algorithm to compress DNA sequences using extended binary trees. J. Theor. Appl. Inform. Tech. 2010, 13, 101–191.

41. Ouyang, J.; Feng, P.; Kang, J. Fast compression of huge DNA sequence data. In Proceedings of the 2012 5th International Conference on BioMedical Engineering and Informatics, Chongqing, China, 16–18 October 2012; pp. 885–888.

42. Li, P.; Wang, S.; Kim, J.; Xiong, H.; Ohno-Machado, L.; Jiang, X. DNA-COMPACT: DNA compression based on a pattern-aware contextual modeling technique. PLoS ONE 2013, 8, e80377. [CrossRef]

43. Roy, S.; Bhagot, A.; Sharma, K.A.; Khatua, S. SBVRLDNAComp: An Effective DNA Sequence Compression Algorithm. Int. J. Comput. Sci. Appl. 2015, 5, 73–85.

44. Roy, S.; Mondal, S.; Khatua, S.; Biswas, M. An Efficient Compression Algorithm for Forthcoming New Species. Int. J. Hybrid Inf. Tech. 2015, 8, 323–332. [CrossRef]

45. Eric, P.V.; Gopalakrishnan, G.; Karunakaran, M. An optimal seed-based compression algorithm for DNA sequences. Adv. Bioinform. 2016. [CrossRef] [PubMed]

46. Rexline, S.J.; Aju, R.G.; Trujilla, L.F. Higher compression from burrows-wheeler transform for DNA sequence. Int. J. Comput. Appl. 2017, 173, 11–15.

47. Keerthy, A.; Priya, S.M. Lempel-Ziv-Welch Compression of DNA Sequence Data with Indexed Multiple Dictionaries. Int. J. Appl. Eng. Res. 2017, 12, 5610–5615.

48. Habib, N.; Ahmed, K.; Jabin, I.; Rahman, M.M. Modified Huﬀbit Compress Algorithm–An Application of R. J. Integr. Bioinform. 2018. [CrossRef] [PubMed]

49. Chen, M.; Li, R.; Yang, L. Optimized Context Weighting for the Compression of the Un-repetitive Genome Sequence Fragment. Wirel. Personal Commun. 2018, 103, 921–939. [CrossRef]

50. Mansouri, D.; Yuan, X. DNA Compression Algorithm. In Proceedings of the International Conference on Neural Information Processing, Siam reap, Cambodia, 13–16 December 2018; Springer: Siam reap, Cambodia; pp. 378–386.

51. Priyanka, M.; Goel, S. A compression algorithm for DNA that uses ASCII values. In Proceedings of the 2014 IEEE International Advance Computing Conference, Gurgaon, India, 21–22 February 2014; pp. 739–743.

52. Bose, T.; Mohammed, M.H.; Dutta, A.; Mande, S.S. BIND–An algorithm for loss-less compression of nucleotide sequence data. J. Biosci. 2012, 37, 785–789. [CrossRef]

53. Jones, D.; Ruzzo, W.; Peng, X.; Katze, M. Compression of next-generation sequencing reads aided by highly efficient de novo assembly. Nucleic Acids Res. 2012, 40. [CrossRef]

54. Uthayakumar, J.; Vengattaraman, T.; Dhavachelvan, P. A new lossless neighborhood indexing sequence (NIS) algorithm for data compression in wireless sensor networks. Ad Hoc Netw. 2019, 83, 149–157.

55. Bakr, N.S.; Sharawi, A.A. Improve the compression of bacterial DNA sequence. In Proceedings of the 2017 13th International Computer Engineering Conference (ICENCO), Cairo, Egypt, 27–28 December 2017; pp. 286–290.

56. National Center for Biotechnology Information. Available online: https://www.ncbi.nlm.nih.gov/ (accessed on 30 March 2019).

57. Roy, S.; Khatua, S. DNA data compression algorithms based on redundancy. Int. J. Found. Comput. Sci. Technol. 2014, 4, 49–58. [CrossRef]

58. Willems, F.M.J.; Shtarkov, Y.M.; Tjalkens, T.J. The context tree weighting method: Basic properties. IEEE Trans. Inf. Theory 1995, 41, 653–664. [CrossRef]