Iterative Spaced Seed Hashing: Closing the gap between spaced seed hashing and k-mer hashing^{*}

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Abstract

Alignment-free classification of sequences has enabled high-throughput processing of sequencing data in many bioinformatics pipelines. Much work has been done to speed-up the indexing of k-mers through hash-table and other data structures. These efforts have led to very fast indexes, but because they are k-mer based, they often lack sensitivity due to sequencing errors or polymorphisms. Spaced seeds are a special type of pattern that accounts for errors or mutations. They allow to improve the sensitivity and they are now routinely used instead of k-mers in many applications. The major drawback of spaced seeds is that they cannot be efficiently hashed and thus their usage increases substantially the computational time.

In this article we address the problem of efficient spaced seed hashing. We propose an iterative algorithm that combines multiple spaced seed hashes by exploiting the similarity of adjacent hash values in order to efficiently compute the next hash. We report a series of experiments on HTS reads

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hashing, with several spaced seeds. Our algorithm can compute the hashing values of spaced seeds with a speedup in range of [3.5x- 7x], outperforming previous methods. Software and Datasets are available at Iterative Spaced Seed Hashing.

Keywords: efficient hashing k-mers spaced seeds

1 Introduction

In computational biology, sequence classification is a common task with many applications such as phylogeny reconstruction (Leimeister et al., 2014), protein classification (Onodera and Shibuya, 2013), metagenomic (Girotto et al., 2016; Marchiori and Comin, 2017; Ounit and Lonardi, 2016). Even if sequence classification is addressable via alignment, the scale of modern datasets has stimulated the development of faster alignment-free similarity methods (Apostolico et al., 2016; Comin and Verzotto, 2014; Comin et al., 2015; Leimeister et al., 2014; Wood and Salzberg, 2014).

The most common alignment-free indexing methods are k-mer based. Large-scale sequence analysis often relies on cataloguing or counting consecutive k-mers (substring of length k) in DNA sequences for indexing, querying and similarity searching. A common step is to break a reference sequence into k-mers and indexing them. An efficient way of implementing this operation is through the use of hash based data structures, e.g. hash tables. Then, to classify sequences are also broken into k-mers and queried against the hash table to check for shared k-mers.

In (Ma et al., 2002) it has been shown that requiring the matches to be nonconsecutive increases the chance of finding similarities and they introduced spaced seeds. They are a modification to the standard k-mer where some positions on the k-mer are set to be don't care, or wildcards, to catch the spaced matches between sequences. In spaced seeds, the matches are distributed so as to maximize the sensitivity, that is the probability to find a local similarity.

Spaced seeds are widely used for approximate sequence matching in bioinformatics and they have been increasingly applied to improve the sensitivity and specificity of homology search algorithms (Kucherov et al., 2006; Noé and Martin, 2014). Spaced seeds are now routinely used, instead of k-mers, in many problems involving sequence comparison like: multiple sequence alignment (Darling et al., 2006), protein classification (Onodera and Shibuya, 2013), read mapping (Rumble et al., 2009), phylogeny reconstruction (Leimeister et al., 2014), metagenome reads clustering and classification (Brinda et al., 2015; Girotto et al., 2017c; Ounit and Lonardi, 2016).

In all these applications, the use of spaced seeds, as opposed to k-mers, has been reported to improve the performance in terms of sensitivity and specificity. However, the major drawback is that the computational cost increases. For example, when k-mers are replaced by spaced seeds, the metagenomic classification of reads of Clark-S (Ounit and Lonardi, 2016) increases the quality of classification, but it also produces a slowdown of 17x with respect to the non-seed version. A similar reduction in time performance when using spaced seeds is reported also in other applications (Brinda et al., 2015; Onodera and Shibuya, 2013; Rumble et al., 2009).

The main reason is that k-mers can be efficiently hashed. In fact, the hashing of a k-mer can be easily computed from the hashing of its predecessor, since they share k-1 symbols. For this reason, indexing all consecutive k-mers in a string can be a very efficient process. However, when using spaced seeds these observations do not longer hold. Therefore, improving the performance of spaced seed hashing algorithms would have a great impact on a wide range of bioinformatics applications. The first attempt to address this question was in the Thesis of R. Harris (Harris, 2007), but hard coding was used to speed-up a non linear packing. Recently, we develop an algorithm based on the indexing of small blocks of runs of matching positions that can be combined to obtain the hashing of spaced-seeds (Girotto et al., 2018a). In (Girotto et al., 2017a, 2018b) we proposed a more promising direction, based on spaced seed self-correlation, in order to reuse part of the hashes already computed. We showed how the hash at position i can be computed based on one best previous hash. Despite the improvement in terms of speedup, the number of symbols that need to be encoded in order to complete the hash could still be high. In this article¹ we solved this problem through: 1) a better way to use previous hashes, maximizing re-use; 2) an iterative algorithm that combines multiple previous hashes. In fact, our algorithm arranges multiple previous hashes in order to recover all k-1 symbols of a spaced seed, so that we only need to encode the new symbol, like with k-mer hashing.

¹A preliminary version of this work was presented at ISBRA 2019(Petrucci et al., 2019)

2 Methods: Iterative Spaced Seed Hashing

2.1 Spaced Seed Hashing: Background

A spaced-seed Q (or just a seed) is a string over the alphabet $\{1, 0\}$ where the 1s correspond to matching positions and 0 to non-matching positions or wildcards, e.g. 1011001. A spaced seed Q can be represented as a set of non negative integers corresponding to the matching positions (1s) in the seed, e.g. $Q = \{0, 2, 3, 6\}$, a notation introduced in (Keich et al., 2004). The weight of a seed, denoted as |Q|, corresponds to the number of 1s, while the length, or span s(Q), is equal to $\max(Q) + 1$.

Given a string x, the positioned spaced seed x[i+Q] identifies a string of length |Q|, where $0 \le i \le n-s(Q)$. The positioned spaced seed x[i+Q], also called Q-gram, is defined as the string $x[i+Q] = \{x_{i+k}, k \in Q\}$.

Example 2.1. Given the seed 1011001, defined as $Q = \{0, 2, 3, 6\}$, with weight |Q| = 4 and span s(Q) = 7. Let us consider the string x = AATCACTTG.

The Q-gram at position 0 of x is defined as x[0+Q] = ATCT. Similarly the other Q-grams are x[1+Q] = ACAT, and x[2+Q] = TACG.

In this article, for ease of discussion, we will consider as hashing function the simple encoding of a string, that is a special case of the Rabin-Karp rolling hash. Later, we will shown how more advanced hashing function can be implemented at no extra cost. Let's consider a coding function from the DNA alphabet $\mathcal{A} = \{A, C, G, T\}$ to a binary codeword, $encode : \mathcal{A} \to \{0, 1\}^{log_2|\mathcal{A}|}$, where encode(A) = 00, encode(C) = 01, encode(G) = 10, and encode(T) = 11. Following the above example, we can compute the encodings of all symbols of the Q-gram x[0 + Q] as follows:

$$\begin{array}{cccc} x[0+Q] & \mathrm{A} & \mathrm{T} & \mathrm{C} & \mathrm{T} \\ encodings & 00 & 11 & 01 & 11 \end{array}$$

Finally, the hashing value of the Q-gram ATCT is 11011100, that is the merge of the encodings of all symbols using little-endian notation. More formally, a standard approach to compute the hashing value of a Q-gram at position i of the string x is the following function h(x[i+Q]):

$$h(x[i+Q]) = \bigvee_{k \in Q} (encode(x_{i+k}) \ll m(k) * log_2|\mathcal{A}|)$$
(1)

Where m(k) is the number of matching positions that appears to the left of k. The function m is defined as $m(k) = |\{i \in Q, \text{ such that } i < k\}|$. In other words, given a position k in the seed, m stores the number of shifts that we need to apply to the encoding of the k-th symbols in order to place it into the hashing. The vector m is important for the computation of the hashing value of a Q-gram.

Example 2.2. In this example, we report an example of hashing value computation for the Q-gram x[1+Q].

Х	А	А	Т	С	А	\mathbf{C}	Т	Т	G
\mathbf{Q}		1	0	1	1	0	0	1	
m		0	1	1	2	3	3	3	
shifted encodings		$\begin{array}{c} 00 \ll 0\\ \underline{00} \end{array}$		01≪2	00≪4			11≪6	
				<u>01</u> 00					
					<u>00</u> 0100				

hashing value

11000100

The previous example shows how the hashing value of x(1+Q) can be computed through the function h(x[1+Q]) = h(ACAT) = 11000100. The hashing value of the other Q-gram can be determined with a similar procedure, i.e. h(x[2+Q]) = h(TACG) = 10010011. The hashing function $h(\cdot)$ is a special case of the Rabin-Karp rolling hash. However, more advanced hashing functions can be defined in a similar way. For example, the cyclic polynomial rolling hash can be computed by replacing: shifts with rotations, OR with XOR, and the function encode() with a table, where DNA characters are mapped to random integers.

In this article we want to address the following problem.

Problem 2.1. Let us consider a string $x = x_0 x_1 \dots x_i \dots x_{n-1}$, of length n, a spaced seed Q and a hash function h that maps strings into a binary codeword. We want to compute all hashing values $\mathcal{H}(x, Q)$ for all the Q-grams of x, starting from the first position 0 of x to the last n - s(Q).

$$\mathcal{H}(x,Q) = \langle h(x[0+Q]), h(x[1+Q]), \dots h(x[n-s(Q)]) \rangle$$

To compute the hash of a contiguous k-mer it is possible to use the hash of its predecessor. In fact, given the hashing value at position i, the hashing for position i + 1 can be obtained with two operations, a shift and the insertion of the encoding of the new symbol, since the two hashes share k - 1 symbols. However, if we consider the case of a spaced seed Q, we can clearly see that this observation does not hold. In fact, in the above example, two consecutive Q-grams, like x[0 + Q] = ATCT and x[1 + Q] = ACAT, do not necessarily have much in common. Since the hashing values are computed in order, the idea is to speed up the computation of the hash at a position i by reusing part of the hashes already computed at previous positions. In this paper we present a solution for Problem 2.1 that maximizes the re-use of previous hashes so that only one symbol needs to be encoded in the new hash, as with k-mers hashing.

2.2 Iterative Spaced Seed Hashing

In the case of spaced seeds, one can reuse part of previous hashes to compute the next one, however we need to explore not only the hash at the previous position, as with k-mers, but the s(Q) - 1 previous hashes. A first attempt to solve this problem was recently proposed in (Girotto et al., 2018b), where the hash at position *i* is computed based on one best previous hash. Despite the improvement in terms of speedup with respect to the standard hashing method, the number of symbols that need to be read in order to complete the hash could still be high. In this article we reduced this value to just one symbol by working in two directions: 1) we devise a better way to use a previous hash, maximizing re-use 2) we propose an iterative algorithm that combines multiple previous hashes.

Let us assume that we want to compute the hashing value at position i and that we already know the hashing value at position i - j, with j < s(Q). We can introduce the following definition of $C_{g,j} = \{k \in Q : k + j \in Q \land m(k) = m(k+j) - m(j) + m(g)\}$ as the positions in Q that after j shifts are still in Q with the propriety that k and k + j positions are both in Q and they are separated by j - g - 1 (not necessarily consecutive) ones. In other words if we are processing the position i of x and we want to reuse the hashing value already computed at position i - j, $C_{g,j}$ represents the symbols, starting at position g of h(x[i - j + Q]), that we can keep while computing h(x[i + Q]).

Example 2.3. Let's consider $Q = \{0, 1, 2, 4, 6, 8, 10\}$. If we know the first

hashing value h(x[0+Q]) and we want to compute the second hash h(x[1+Q]), the following example show how to construct $C_{0,1}$.

0	1					0,1						
k		0	1	2	3	4	5	6	$\overline{7}$	8	9	10
Q		1	1	1	0	1	0	1	0	1	0	1
$Q \ll 1$	1	1	1	0	1	0	1	0	1	0	1	
m(k)		0	1	2	3	3	4	4	5	5	6	6
m(k+1)-m(1)+m(0)	-1	0	1	2	2	3	3	4	4	5	5	
$C_{0,1}$		0	1									

The symbols at positions $C_{0,1} = \{0, 1\}$ of the hash h(x[1+Q]) have already been encoded in the hash h(x[0+Q]) and we can keep them. In order to complete h(x[1+Q]), the number of remaining symbols are $|Q| - |C_{0,1}| = 5$.

In the article (Girotto et al., 2018b) we use only the symbols in $C_{0,j}$, that is g was always 0. As we will see in the next examples, if we are allowed to remove the first g symbols from the hash of h(x[i - j + Q]), we can recover more symbols in order to compute h(x[i + Q]).

Example 2.4. Let us consider the hash at position 2 h(x[2+Q]), and the hash at position 0 h(x[0+Q]). In this case we are interested in $C_{0,2}$.

-	· · · ·	L .	• 1 /							~,	-		
k			0	1	2	3	4	5	6	7	8	9	10
Q			1	1	1	0	1	0	1	0	1	0	1
$Q \ll 2$	1	1	1	0	1	0	1	0	1	0	1		
m(k)			0	1	2	3	3	4	4	5	5	6	6
m(k+2)-m(2)+m(0)	-2	-1	0	1	1	2	2	3	3	4	4		
$C_{0,2}$			0										

Thus, the only position that we can recover is $C_{0,2} = \{0\}$. On the other hand, if we are allowed to skip the first position of the hash h(x[0+Q]) and consider $C_{1,2}$, instead of $C_{0,2}$, we have:

k			0	1	2	3	4	5	6	$\mid 7$	8	9	10
Q			1	1	1	0	1	0	1	0	1	0	1
$Q \ll 2$	1	1	1	0	1	0	1	0	1	0	1		
m(k)			0	1	2	3	3	4	4	5	5	6	6
m(k+2)-m(2)+m(1)	-1	0	1	2	2	3	3	4	4	5	5		
$C_{1,2}$					2		4		6		8		

Where, we can re-use the symbols $C_{1,2} = \{2, 4, 6, 8\}$ of h(x[0+Q]) in order to compute h(x[2+Q]). This example shows how the original definition of C_j in (Girotto et al., 2018b), that in this work corresponds to $C_{0,2} = \{0\}$, was not optimal and more symbols could be recovered from the same hash with $C_{1,2} = \{2, 4, 6, 8\}.$

In (Girotto et al., 2018b), the hash value at a given position was reconstructed starting from the best previous hash. However, the number of symbols to be inserted to complete the hash could still be high. In this article we propose a new method that not only consider the best previous hash, but all previous hashes at once. For a given hash to be computed h_i , we devised an iterative algorithm that is able to find a combination of the previous hashes that covers all symbols of h_i , apart from the last one. That is, we can combine multiple hashes in order to recover |Q| - 1 symbols of h_i , so that we only need to read the new symbol, like with k-mer hashing.

Let us assume that we have already computed a portion of the hash h_i , and that the remaining symbols are $Q' \subset Q$. We can search the best previous hash that covers the largest number of positions of Q'. To this end, we define the function BestPrev(s, Q') that searches for this best previous hash:

$BestPrev(s,Q') = argmax_{z \in [0,s-1],k \in [1,s]} |\mathcal{C}_{z,k} \cap Q'|$

This function will return a pair (g, j) that identifies the best previous hash at position h_{i-j} from which, after removing the first g symbols, we can recover $|C_{g,j} \cap Q'|$ symbols. In order to extract these symbols from h_{i-j} we define a mask, $Mask_{g,j}$, that filters these positions. The algorithm iteratively searches the best previous hashes, until all |Q| - 1 symbols have been recovered. An overview of the method is shown below:

Our iterative algorithm scans the input string x and computes all hashing values according to the spaced seed Q. In order to better understand the amount of savings we evaluate the algorithm by counting the number of symbols that are read and encoded. First, we can consider the input string to be long enough so that we can discard the transient of the first s(Q)-1 hashes. Let us continue to analyze the spaced seed 11101010101, that corresponds to $Q = \{0, 1, 2, 4, 6, 8, 10\}$. If we use the standard function h(x[i + Q]) to compute all hashes, each symbol of x is read |Q| = 7 times.

Algorithm 1 Iterative Spaced Seed Hashing

1: Compute $C_{q,k}$ and $Mask(g,k) \forall g,k$; 2: $h_0 := \text{compute } h(x[0+Q]);$ 3: for i := 1 to s(Q) - 1 do Q' = Q;4: while $|Q'| \neq 1$ do 5:(g,k) = BestPrev(i,Q');6: if $(Q' \cap C_{q,k}) == \emptyset$ then 7: Exit while; 8: else 9: $h_i := h_i \text{ OR } ((h_{i-k} \text{ AND } Mask(g,k)) >> k * log_2|\mathcal{A}|);$ 10: $Q' = Q' - C_{q,k} ;$ 11: end if 12:end while 13:for all $k \in Q'$ do 14:insert $encode(x_{i+k})$ at position $m(k) * log_2|\mathcal{A}|$ of h_i ; 15:end for 16:17: end for 18: for i := s(Q) to |x| - s(Q) do Q' = Q;19:while $|Q'| \neq 1$ do 20: (g,k) = BestPrev(s(Q) - 1, Q');21: $h_i := h_i \text{ OR } ((h_{i-k} \text{ AND } Mask(g, k)) >> k * log_2|\mathcal{A}|);$ 22: $Q' = Q' - C_{q,k} ;$ 23:end while 24: insert $encode(x_{i+s(Q)-1})$ at last position of h_i ; 25:26: end for

In the first iteration of our algorithm (lines=19-25) Q' = Q and the best previous hash BestPrev(s(Q) - 1, Q') = (1, 2) is $C_{1,2} = \{2, 4, 6, 8\}$. Thus, while computing h_i we can recover these 4 symbols from h_{i-2} . At the end of the first iteration Q' is updated to $\{0, 1, 10\}$. During the second iteration the best previous hash BestPrev(s(Q) - 1, Q') = (0, 1) is $C_{0,1} = \{0, 1\}$. As above, we can append these two symbols from h_{i-1} to the hash h_i . Now, we have that $Q' = \{10\}$, that is only one symbol is left. The last symbol is read and encoded into h_i , and the hash is complete. In summary, after two iterations all |Q| - 1 symbols of h_i have been encoded into the hash, and we only need to read one new symbol from the sequence. Moreover, if one needs to scan a string with a spaced seed and to compute all hashing values, the above algorithm guarantees to minimize the number of symbols to read. In fact, with our algorithm, we can compute all hashing values while reading each symbol of the input string only once, as with k-mers.

3 Results and discussion

In this section we will present the results of some experiments in which ISSH is compared against two other approaches available in literature: FISH (Girotto et al., 2018a) (block-based) and FSH (Girotto et al., 2018b) (overlap-based).

3.1 Experimental settings

We use the same settings as in previous studies (Girotto et al., 2018a,b). The spaced seeds belong to three different types of spaced seeds, according to the objective function used to generate them: maximizing the hit probability (Ounit and Lonardi, 2016); minimizing the overlap complexity (Hahn et al., 2016); and maximizing the sensitivity (Hahn et al., 2016). We tested three spaced seeds for each type, all with weight W = 22 and length L = 31. This list of spaced seeds is presented in Table of 1 with labels from Q1 to Q9. Furthermore, we used other sets of spaced seeds, built with *rashbari* (Hahn et al., 2016), which have weights from 11 to 32 and the same length. The complete list of the spaced seeds used is reported in the Appendix (see Tables 3-5). The datasets of metagenomic reads to be hashed were taken from previous papers on binning and classification (Girotto et al., 2016, 2017b; Wood and Salzberg, 2014). Details about the reads datasets are shown in Table 2 in the Appendix. All the experiments have been performed on a laptop equipped with an Intel i7-3537U CPU at 2 GHz and 8 GB of RAM.

3.2 Analysis of the Time Performances

The first comparison we present is between the performances of ISSH, FISH and FSH in terms of speedup with respect to the standard hash computation (i.e. applying Eq.1 to each position). Figure 1 shows the average speedup among all datasets, for each of the spaced seeds Q1-Q9, obtained by the three different methods.

Table 1: The spaced seeds Q1-Q9 deployed in the experiments grouped by their type.

Space	ced seeds maximizing the hit probability Ounit and Lonardi (2016)
Q1	11110111011100101110010110111111
Q2	11111010111001011011100110111111
Q3	11111010011101011011001110111111
Sp	baced seeds minimizing the overlap complexity Hahn et al. (2016)
Q4	1111010111010011001110111110111
Q5	11101110111011110100101100111111
Q6	11111010010111001111101011011111
	Spaced seeds maximizing the sensitivity Hahn et al. (2016)
Q7	1111011110011010111110101011011
Q8	11101010111011001101001111111111
Q9	1111110101101011100111001111



Figure 1: The average speedup obtaind by ISSH, FISH and FSH with respect to the standard computation.

It can be seen that ISSH is much faster than both FISH and FSH for all the spaced seeds. In terms of actual running time, the standard approach (Eq.1) requires about 14 minutes to compute the hashes for a single spaced seed on all datasets. ISSH takes just over 2 minutes with an average speedup of 6.2. As for the other two approaches, FISH and FSH, they compute the hashes in 6 and 9 minutes respectively, with an average speedup of 2 (FISH) and 1.5 (FSH).

We also notice that the variation among the speedups, relative to different spaced seeds using the same method, are lower for ISSH, for which the speedups are in the range [6.05-6.25] while for FISH and FSH the range is [1.89-2.16] and [1.18-1.58], respectively. For all the tested methods there is a correlation between the spaced seed structure and the time needed for the computation. FISH depends on the number of blocks of 1s, while both ISSH and FSH depend on the spaced seed self-correlation. ISSH performances are also sensitive to the number of iterations. However, the experiments show that, even if FSH performs a single iteration, the time required to naively compute the hash for all the non-overlapping positions is more than the time required by ISSH to perform more iterations. Moreover, for all the tested spaced seeds the number of iterations needed by ISSH was on average 4.

Figure 2 gives an insight on the performance of ISSH with respect to each spaced seed and each datasets considered.



Figure 2: Speedup of ISSH of all the single spaced seeds for each of the considered datasets, ordered by reads length.

First of all, we notice that the performances are basically independent on the spaced seed used. Next, for what concerns the datasets characteristics, it can be observed that the speedup increases with the reads length, reaching the highest values for the datasets R7, R8 and R9, which have the longest reads. This behavior is expected: when considering longer reads the slowdown caused by the initial transient – in which more than one symbol has to be encoded – is less relevant with respect to the total running time.

In Figure 3 we report the speedups on each datasets obtained by Q7, a typical spaced seed (the other spaced seeds performances are similar) using ISSH, FISH and FSH.



Figure 3: Details of the speedup on the spaced seed Q7 on each datasets, ordered by reads length, using ISSH, FISH and FSH.

All the results are compatible with the above observations: ISSH, if compared to FISH and FSH, allows to compute the hashing values faster for all the datasets. Futhermore, by using ISSH, the improvement on long reads datasets is larger than the improvement obtained with FISH or FSH.

3.3 Effect of Spaced Seeds Weight on Time Performances

The experiments presented here point out the connection between the density of a spaced seed and the speedup. We considered four sets of nine spaced seeds, generated with *rasbhari* (Hahn et al., 2016), with weights 14, 18, 22 and 26 and a fixed length of 31, see Tables 3-5 in the Appendix.

In Figure 4 we compare the average speedup of ISSH, FISH and FSH for these sets of spaced seeds as a function of the weight W. We notice that the speedup grows as the weight increases. This phenomenon is consistent among all the methods we analyzed. It is reasonable to think that such difference is due to how the hashes are computed with the standard method using Eq.1



Figure 4: The speedup of ISSH, FISH and FSH as a function of the spaced seeds density (L=31 and W=14, 18, 22, and 26).

(against which all methods are compared), because denser spaced seeds imply hashes with a larger number of symbols that need to be encoded and joined together. Moreover, for ISSH we have that denser spaced seeds have more chances of needing fewer previously calculated hashes to compute each of the |Q| - 1 symbols, thus saving further iterations.

Both these effects are emphasized when looking at the actual running times needed by the least dense group (W = 14) and by the most dense group (W = 26) of spaced seeds. The standard method requires 9.73 and 15.11 minutes, respectively, while ISSH spends only 2.75 and 2.16 minutes to perform the same task.

3.4 Effect of Number of Iterations on Time Performances

The experiments described in this section have been essential to understand how many of the previously calculated hashes we should use in order to further speed up the computations. At the beginning we were not sure whether it was better or not to increase the number of iterations, for each of the |x| - s(Q)hashes that need to be computed, to recover only few symbols. To address this problem we considered the speedup obtained with ISSH by progressively limiting the number of iterations, that is the number of previous hashes used to compute a new one, from one to five (number needed to cover all the |Q|-1 symbols for all the spaced seeds considered). The results are shown in Figure 5 where the spaced seeds used have length L = 31 and weight W = 14.



Figure 5: Speedup of ISSH, on the seeds of table 3 (see Appendix), with an increasing limit to the number of iterations used to compute each hash.

To compute the remaining symbols it has been used a similar approach to the one described in Girotto et al. (2018b) for FSH, which is also used by ISSH to complete the hashes computed in the transient part. We can see that gradually increasing the number of iterations the speedup becomes greater: from speedups similar to the ones obtained with FSH, which only considers one single previous hash (that correspond to a single ISSH iteration), to the higher speedups of this new approach.

Here we can also observe again that the number of iterations has an impact on the speedup. For the spaced seed Q18 only 4 iterations are required to recover all the |Q| - 1 positions, one less than the other spaced seeds, and this leads to a higher speedup for Q18 if compared to the other spaced seeds with the same density.

In Figure 6 the same speedups shown in Figure 5 are plotted, but this time against the number of symbols recovered.

Even in this case, it is clear that as ISSH recovers more symbols – which means having to apply the encode function for fewer positions – the speedup increases. Another interesting observation is that a small variation is present between the speedup of computations in which the same number of symbols are recovered, but the number of iterations used is different. For example, if we consider the cases in which 10 symbols are recovered, the speedups are



Figure 6: Speedup of ISSH obtained recovering a certain number of symbols for the seeds of table 3 (see Appendix).

almost the same, but for Q14 and Q15 two iterations are needed while for Q10 an additional one is required. In summary, the speedup depends more on the number of symbols recovered rather than the number of iterations.

4 Conclusions

In this article we present ISSH (Iterative Spaced Seed Hashing), an iterative algorithm that combines multiple previous hashes in order to maximize the re-use of already computed hash values. The average speedup of ISSH with respect to the standard computation of hash values is in range of [3.5x- 7x], depending on spaced seed density and reads length. In all experiments ISSH outperforms previously proposed algorithms. Possible directions of research are the combination of multiple spaced seeds and the investigation of global optimization schemes.

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5 Appendix

Da	atasets	Number of reads	Avg. read length
	S6	1426457	80
	S7	3307100	80
	S9	4468336	80
	S10	9981172	80
	L5	1016418	80
	L6	1182178	80
I	HiSeq	9989713	91
si	$\mathrm{mBA5}$	1631100	100
Ν	/ixK1	9629886	101
Ν	/ixK2	7149900	101
Ν	MiSeq	3990282	131
	R7	290473	702
	R8	374576	715
	R9	588256	715

Table 2: Number of reads and average lengths for each of the dataset used in our experiments.

Table 3: Nine spaced seeds with W = 14 and length 31 computed with rashhari minimizing overlap complexity.

Q10	1110000100110100110110001010001
Q11	1011000001001011111101000100001
Q12	1010000001001101100110000110111
Q13	1110010001000010000111011100011
Q14	1010000111100111100011000000011
Q15	10001000001000001010111111011
Q16	1011100000100101011100010110001
Q17	1000101011100101101010001100001
Q18	1011010110001010101000000010111

Table 4: Nine spaced seeds with W = 18 and length 31 computed with rashari minimizing overlap complexity.

1011010110010101111011010110001
1011001001011011111101010101001
1010101001011101100110000111111
1110110011000010001111011110011
1010011111101111101011000000011
110010010101000011010111111011
1011100100110101111100010111001
1110101011100111101010011100001
1011011110001011101000100011111

Table 5: Nine spaced seeds with W = 26 and length 31 computed with rashbari minimizing overlap complexity.

0	1 1 7	
Q28	1111101110111011111101101111111	
Q29	11111110111111111111111010101101	
Q30	1110111011011111111110011111111	
Q31	1111111111100111101111111110011	
Q32	1111011111111111010110111110111	
Q33	11011011011111001111111111111111	
Q34	10111101011111011111111111111111	
Q35	1111101111110111111110111110011	
Q36	1011111111101011111010111111111	