

Analytic combinatorics for bioinformatics I: seeding methods

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Abstract

The abstract will come later.

1 Introduction

High throughput sequencing is changing the face of biology [7,14]. More data is of course better, but recently, technical improvements have started to outpace the progress of algorithms [13]. When the problems are too large, one has to replace exact algorithms by heuristics that are much faster, but do not guarantee to return the right result. Good heuristics are all about understanding the input data. With the right data model, we can calculate the risk of not returning the right answer and adjust the algorithm to achieve more precision or more speed. When the data is poorly understood, heuristics may be slow or inefficient for unknown reasons.

A particular area of bioinformatics where heuristics have been in use for a long time is the field of sequence alignment [12]. Computing the best alignment between two sequences is carried out by dynamic programming in time $O(mn)$, where m and n are the sequence lengths [2]. When at least one of the sequences is long (*e.g.* a genome), this is prohibitive and heuristics are required.

The most studied heuristics for sequence alignment are called seeding methods [15]. In a nutshell, the idea is to search short regions of the two sequences that are very similar and use them as candidates to anchor the dynamic programming alignment, which is performed only locally *i.e.* between subsequences of the input. These short regions of high similarity are called “seeds”. The benefit of the approach is that seeds can be found in short time. The risk is that they may not exist, even if the input sequences are similar.

This strategy was most famously implemented in BLAST for the purpose of finding local homology between proteins [1]. By working out an approximate distribution of the identity score for the seeds [8, 9], the authors were able to calibrate the BLAST heuristic very accurately in order to gain speed. The

algorithm always performs the minimum amount of work for a desired confidence level.

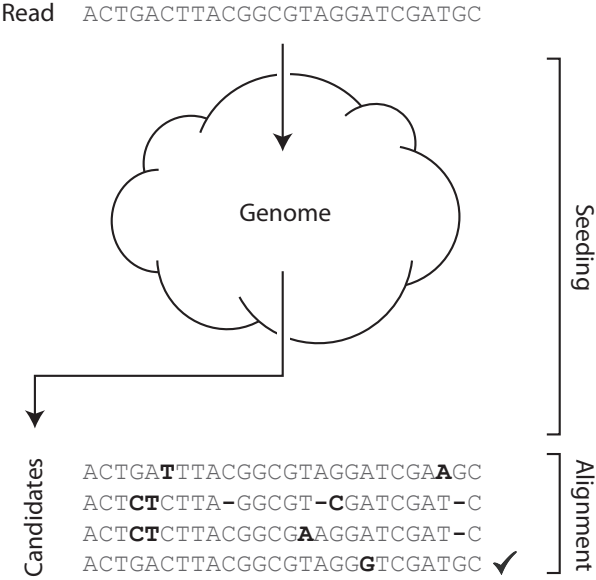


Figure 1: Seeding heuristic in the mapping problem. The sequencing read belongs to an unknown portion of the genome; the task is to find its original sequence (and usually its location). Because of possible read errors, the read may not be identical to the original sequence. Most algorithms feature a seeding stage and an alignment stage. The purpose of seeding is to output a short list of candidate sequences, that of alignment is to determine which is the best. Seeding is a heuristic because it does not guarantee that the overall best sequence is in the list of candidates.

Seeding methods are also heavily used in the mapping problem (see figure 1), where the original sequence of a read must be found in a reference genome. The discovery of indexing methods based on the Burrows-Wheeler transform [3] was instrumental to develop short read mappers such as BWA and Bowtie [10, 11]. With such indexes, one can know the number of occurrences of a substring in a genome in time independent of the genome size. This yields a powerful seeding strategy whereby all the substrings of the read are queried in the genome. The main limitation is that the seeds must have exactly the same sequence in the read as in the genome.

The heuristic should be calibrated from the probability that a seed of given length can be found in the read, but this problem has not been fully solved. The answer depends on the types and frequencies of errors, which are often context-dependent. Overall, the lack of theoretical framework to model seeding probabilities is slowing progress on this line of research.

Here we solve this problem for arbitrary error models using the powerful

theory of analytic combinatorics. This field of research was most significantly developed by Andrew Odlyzko, Robert Sedgewick and Philippe Flajolet [4–6]. The theory is now mature and used to tackle many problems outside the analysis of algorithms. However, it has not yet been fully realized how useful it can be for bioinformatics.

This document is predominantly written for bioinformaticians and people with a working knowledge of sequencing technologies and their applications. Accordingly, the focus will be on explaining the mathematical concepts, rather than the technological aspects. Also, our goal here is not to push the boundaries of analytic combinatorics, but to explain how its simplest concepts are useful to solve common problems in bioinformatics. We have opted for simplicity, to the detriment of generality and rigor. The results presented here are only a basic introduction to analytic combinatorics; the field is currently much more advanced and we refer the interested to the original literature.

2 Elements of analytic combinatorics

2.1 Weighted generating functions

Definition 1. Let \mathcal{A} be a set of combinatorial objects characterized by a size and a weight. The **weighted generating function** of \mathcal{A} is defined as

$$A(z) = \sum_{a \in \mathcal{A}} w(a) z^{|a|}, \quad (1)$$

where $|a|$ and $w(a)$ denote the **size** and **weight** of the object a , respectively. Expression (1) also defines a sequence $(a_k)_{k \geq 0}$ such that

$$A(z) = \sum_{k=0}^{\infty} a_k z^k. \quad (2)$$

By definition $a_k = \sum_{a \in A_k} w(a)$, where A_k is the class of objects of size k in \mathcal{A} . The number a_k is called the total weight of objects of size k .

Remark 1. Expression (2) shows that the terms a_k are the coefficients of the Taylor series of $A(z)$.

Remark 2. If the weight of every object $a \in \mathcal{A}$ is 1, then $A(z)$ is called a simple generating function, and in expression (2) $a_k = |A_k|$ is the number of objects of size k in \mathcal{A} .

Remark 3. Typical combinatorial objects include binary trees, permutations, derangements, multisets etc., but here we will focus exclusively on sequences of letters from finite alphabets. In what follows, the reader can think of “objects” as “finite sequences of letters”.

Expressions (1) and (2) are of course equivalent. Depending on the context, we will use one or the other.

Example 1. Let $\mathcal{A} = \{a\}$ and $\mathcal{B} = \{b\}$ be alphabets with a single letter (of size 1). Assume $w(a) = p$ and $w(b) = q$. The weighted generating functions of \mathcal{A} and \mathcal{B} are then $A(z) = pz$ and $B(z) = qz$, respectively.

*

The motivation for definition 1 is that operations on combinatorial objects translate into operations on their weighted generating functions. With the methods detailed below, we will obtain the weighted generating functions of elaborate combinatorial objects from simple ones. Using equation (2), we will extract the weight or probability of objects of size k from such weighted generating functions.

This approach is counter-intuitive at first, but propositions 4 and 5 below will show how we can compute probabilities that are inaccessible to more frontal approaches.

Let us start with the most straightforward ways to obtain new weighted generating functions: additions and multiplications. If $A(z)$ and $B(z)$ are the weighted generating functions of two mutually exclusive sets \mathcal{A} and \mathcal{B} , the weighted generating function of $\mathcal{A} \cup \mathcal{B}$ is $A(z) + B(z)$, as appears immediately from expression (1).

Example 2. With the definitions of example 1, the weighted generating function of the alphabet $\mathcal{A} \cup \mathcal{B} = \{a, b\}$ is $pz + qz = A(z) + B(z)$.

*

Size and weight can be defined for pairs of objects in $\mathcal{A} \times \mathcal{B}$ as $|(a, b)| = |a| + |b|$ and $w(a, b) = w(a)w(b)$. In other words the sizes are added and the weights are multiplied. With this convention, the weighted generating function of the Cartesian product $\mathcal{A} \times \mathcal{B}$ is $A(z)B(z)$. This simply follows from expression (1) and

$$A(z)B(z) = \sum_{a \in \mathcal{A}} w(a)z^{|a|} \sum_{b \in \mathcal{B}} w(b)z^{|b|} = \sum_{(a,b) \in \mathcal{A} \times \mathcal{B}} w(a)w(b)z^{|a|+|b|}.$$

Example 3. With the definitions of example 1, $\mathcal{A}^2 = \{(a, a)\}$ contains a single pair of letters, with size 2 and weight p^2 . Its weighted generating function is $p^2z^2 = A(z)^2$.

*

Example 4. Still with the definitions of example 1, $(\mathcal{A} \cup \mathcal{B})^2$ contains the four pairs of letters (a, a) , (a, b) , (b, a) and (b, b) . They have size 2 and respective weight p^2 , pq , qp , and q^2 , so the weighted generating function of $(\mathcal{A} \cup \mathcal{B})^2$ is $(p^2 + 2pq + q^2)z^2 = (A(z) + B(z))^2$.

*

We can further extend the definition of size and weight to any finite Cartesian product in the same way. The sizes are always added and the weights are always multiplied. The generating function of a cartesian product then comes as the product of their generating functions. This allows us to construct finite sequences of objects.

Example 5. With the definitions of example 1, $\mathcal{A}^k = \{(a, a, \dots, a)\}$ contains a single k -tuple of letters, with size k and weight p^k , so its weighted generating function is $p^k z^k$. Since the sets $\mathcal{A}, \mathcal{A}^2, \mathcal{A}^3, \dots$ are mutually exclusive, the weighted generating function of their union is

$$pz + p^2 z^2 + p^3 z^3 + \dots$$

This infinite Taylor series can be expressed as a simple function. For any $k \geq 1$, $(1 - pz)(pz + (pz)^2 + \dots + (pz)^k) = pz - (pz)^{k+1}$. If $|z| < 1/p$, the term $(pz)^{k+1}$ vanishes as k increases. So the weighted generating function of $\cup_{k \geq 1} \mathcal{A}^k$ is defined for $|z| < 1/p$ and is equal to

$$pz + (pz)^2 + (pz)^3 + \dots = \frac{pz}{1 - pz}.$$

*

Example 5 is fundamental. It can be generalized to any set \mathcal{A} of combinatorial objects.

Proposition 1. *Let \mathcal{A} be a set with weighted generating function $A(z)$. The weighted generating function of $\mathcal{A}^+ = \cup_{k=1}^{\infty} \mathcal{A}^k$, called the set of **non-empty sequences** of elements of \mathcal{A} is defined for $|A(z)| < 1$ and is equal to*

$$\frac{A(z)}{1 - A(z)}.$$

Proof. For $k \geq 1$, the weighted generating function of \mathcal{A}^k is $A(z)^k$. Since the sets \mathcal{A}^k are mutually exclusive, the weighted generating function of their union is $A(z) + A(z)^2 + \dots = A(z)/(1 - A(z))$, provided $|A(z)| < 1$. \square

We introduce the **empty object** ε , which has size 0 and weight 1. Its weighted generating function is thus 1. By convention, we define the zeroth power of a combinatorial set \mathcal{A} as $\mathcal{A}^0 = \{\varepsilon\}$. With this definition, we can state a variant of the previous proposition.

Proposition 2. *Let \mathcal{A} be a set with weighted generating function $A(z)$. The weighted generating function of $\mathcal{A}^* = \cup_{k=0}^{\infty} \mathcal{A}^k$, called the set of **sequences** of elements of \mathcal{A} is defined for $|A(z)| < 1$ and is equal to*

$$\frac{1}{1 - A(z)}.$$

Proof. $\mathcal{A}^* = \{\varepsilon\} \cup \mathcal{A}^+$. By proposition 1, the weighted generating function is $1 + A(z)/(1 - A(z)) = 1/(1 - A(z))$, provided $|A(z)| < 1$. \square

Remark 4. These expressions are not defined for $A(z) = 1$, i.e. when \mathcal{A} contains only the empty object. In other words, one cannot construct sequences of empty objects. More generally, we will always assume that the sets used to construct sequences do not contain the empty object ε . Otherwise, ε is present in each \mathcal{A}^k and they are not mutually exclusive.

Remark 5. If $|A(z)| > 1$, the function $A(z)/(1 - A(z))$ is well defined but the series $A(z) + A(z)^2 + \dots$ does not converge. This is not a problem, as relation (2) does not need to hold for every z . However, in what follows we will always assume that it does. In other words, we will always assume that $|z|$ is smaller than the radius of convergence of the weighted generating function of interest.

It is important to insist that the expression “sequences of” refers to the set \mathcal{A}^* and not to \mathcal{A}^+ , i.e. the empty object ε is a sequence of anything. For clarity, we will systematically use the expression “non-empty sequences of” when ε is excluded.

Example 6. Following the definitions of example 1, $(\mathcal{A} \cup \mathcal{B})^*$ is the set of sequences of a 's and b 's. By proposition 2 and example 2, the weighted generating function of this set is

$$\frac{1}{1 - (p + q)z}.$$

The function $1/(1 - (p + q)z)$ is just a compact representation of the infinite series $1, (p + q), (p + q)^2, (p + q)^3, \dots$ It thus carries all the information about the total weights of the sequences of any size. If $p = q = 1$, i.e. if we count the total amount of sequences with simple generating functions (see remark 2), we obtain $1/(1 - 2z) = 1 + 2z + 4z^2 + 8z^3 + 16z^4 + \dots$ meaning that there are 2^k sequences of size k . If p and q are the respective probabilities of occurrence of a and b , with $p + q \leq 1$, then the equality $1/(1 - (p + q)z) = 1 + (p + q)z + (p + q)^2z^2 + (p + q)^3z^3 + \dots$ means that the probability that a sequence of size k contains only a 's and b 's is $(p + q)^k$.

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These simple examples are better solved by intuition. However, we will soon see that analytic combinatorics allows us to solve a wider range of problems.

2.2 Sequences and transfer matrices

In many combinatorial applications, one needs to count the sequences where a pattern does not occur, or where some symbols may not follow each other. A convenient way to find the weighted generating functions of such sequences is to encode this information in so-called “transfer matrices”. We will illustrate the process with an example from biology.

In many animal genomes, DNA methylation can only occur on the dinucleotide **CG**. Because of this property, it is interesting to count the sequences

that have no CG. Sequences without CG can be thought of as walks on a directed graph with restricted transitions. Nucleotides are represented as nodes, and edges indicate that the nucleotide at the tip can follow the nucleotide at the base. Sequences without CG correspond to walks on a complete graph where the edge from C to G is removed (see figure 2).

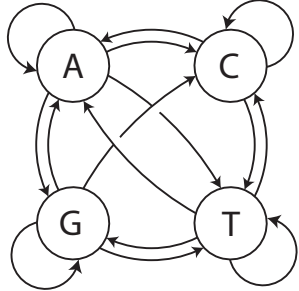


Figure 2: **Graph representation of sequences of nucleotides without CG.** A sequence without CG is equivalent to a walk on this graph. All the transitions are allowed, except C to G.

The same graph can also be represented as an adjacency matrix M , where a 1 at position (i, j) indicates that there is an edge from node i to node j , and a 0 indicates that there is no edge. For instance, the adjacency matrix of the graph of figure 2 is

$$M = \begin{matrix} & \begin{matrix} A & C & G & T \end{matrix} \\ \begin{matrix} A \\ C \\ G \\ T \end{matrix} & \begin{bmatrix} 1 & 1 & 1 & 1 \\ 1 & 1 & 0 & 1 \\ 1 & 1 & 1 & 1 \\ 1 & 1 & 1 & 1 \end{bmatrix} \end{matrix}.$$

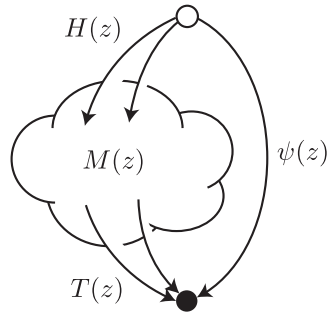
The main advantage of the adjacency matrix representation is that the powers of M give an analytical way to count the walks on the graph. The entry of M^n at position (i, j) is the number of walks of n steps that start with node i and end with node j . So we can count sequences without CG by computing the successive powers of the adjacency matrix M above.

The same idea can be used to find the weighted generating function of sequences of combinatorial objects. We specify the internal structure of sequences by a “transfer graph”, where the edges represent classes of combinatorial objects.

Definition 2. A **transfer graph** is a directed graph whose edges are weighted by generating functions. In addition, a transfer graph must contain a head vertex with only outgoing edges, and a tail vertex with only incoming edges. The graph can be represented as a matrix called a **transfer matrix**.

Following the edges of a transfer graph from the head vertex to the tail vertex describes a sequence of combinatorial objects. The associated weighted generating function is the product of the functions labelling the edges (thus an absent edge is associated to the function 0).

Transfer graphs have the general structure below. The head vertex is represented as an open circle and the tail vertex as a closed circle. The edge labelled $\psi(z)$, from the head vertex to the tail vertex represents additional sequences. Typically, $\psi(z) = 1$, as the only sequence that needs to be added explicitly is the empty object ε .

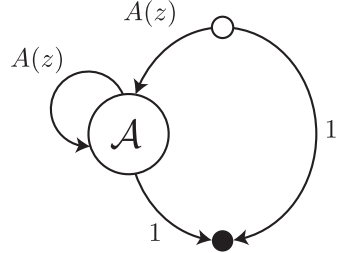


The “body” of the graph, represented as a cloud, contains the remaining m vertices. $M(z)$ is the $m \times m$ matrix representation of the corresponding subgraph. $H(z)$ is a vector of m weighted generating functions associated with the m edges from the head vertex to the body of the graph, and $T(z)$ is a vector of m weighted generating functions associated with the edges from the body of graph to the tail vertex. $H(z)$ and $T(z)$ are called the “head” and “tail” vectors, respectively.

Remark 6. Transfer graphs are reminiscent of deterministic finite automata (DFA). The initial and accept states of a DFA correspond to the head and tail vertices of a transfer graph. The alphabet of a DFA corresponds to the combinatorial objects that can be concatenated (or to their weighted generating functions) and the transition function corresponds to the transfer matrix. We will not make further use of this analogy in this document.

Definition 3. The **sequences generated by a transfer matrix** correspond to the paths from the head vertex to the tail vertex of the associated transfer graph.

Example 7. Let \mathcal{A} be a class of combinatorial objects with weighted generating function $A(z)$. Sequences of objects from \mathcal{A} can be thought of as walks on the graph below.



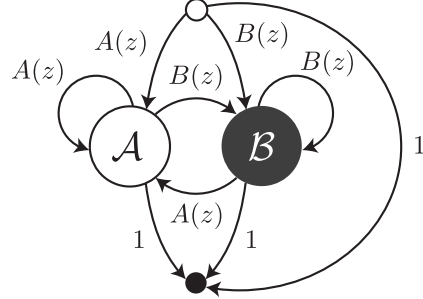
The edge from the vertex \mathcal{A} to the tail vertex marks the end of the sequence, by appending a final empty object ε with weight 1. The transfer matrix of this graph is the 3×3 matrix

$$M_*(z) = \begin{matrix} & \circ & \mathcal{A} & \bullet \\ \circ & 0 & A(z) & 1 \\ \mathcal{A} & 0 & A(z) & 1 \\ \bullet & 0 & 0 & 0 \end{matrix}.$$

The head vector $H(z)$ has dimension 1 and is equal to $A(z)$. The tail vector $T(z)$ also has dimension 1 and is equal to 1. $M(z)$, the matrix of the body of the graph is the 1×1 matrix $[A(z)]$.

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Example 8. Let \mathcal{A} and \mathcal{B} be mutually exclusive classes of combinatorial objects with respective weighted generating function $A(z)$ and $B(z)$. Sequences of objects from \mathcal{A} or \mathcal{B} can be thought of as walks on the graph below.



The transfer matrix of this graph is the 4×4 matrix

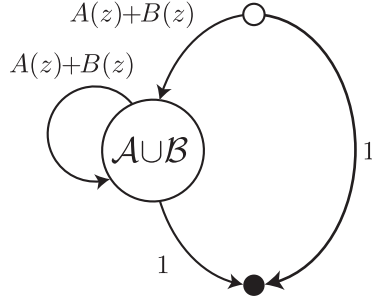
$$M_*(z) = \begin{matrix} & \circ & \mathcal{A} & \mathcal{B} & \bullet \\ \circ & 0 & A(z) & B(z) & 1 \\ \mathcal{A} & 0 & A(z) & B(z) & 1 \\ \mathcal{B} & 0 & A(z) & B(z) & 1 \\ \bullet & 0 & 0 & 0 & 0 \end{matrix}.$$

The head vector $H(z)$ has dimension 2 and is equal to $(A(z), B(z))^\top$. The tail vector $T(z)$ also has dimension 2 and is equal to $(1, 1)^\top$. $M(z)$, the matrix of the body of the graph is the 2×2 matrix

$$M(z) = \begin{matrix} & \mathcal{A} & \mathcal{B} \\ \mathcal{A} & \begin{bmatrix} A(z) & B(z) \\ A(z) & B(z) \end{bmatrix} \\ \mathcal{B} & & \end{matrix}.$$

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Example 9. With the assumptions of example 8, sequences of objects from \mathcal{A} or \mathcal{B} can also be thought of as walks on a the graph below.



Indeed, because \mathcal{A} and \mathcal{B} are mutually exclusive, we know that the weighted generating function of $\mathcal{A} \cup \mathcal{B}$ is $A(z) + B(z)$. The transfer matrix of this graph is the 3×3 matrix

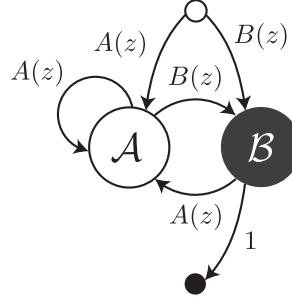
$$M_*(z) = \begin{matrix} & \circ & \mathcal{A} \cup \mathcal{B} & \bullet \\ \circ & \begin{bmatrix} 0 & A(z) + B(z) & 1 \\ 0 & A(z) + B(z) & 1 \\ 0 & 0 & 0 \end{bmatrix} \\ \mathcal{A} \cup \mathcal{B} & & & \\ \bullet & & & \end{matrix}.$$

The head vector $H(z)$ has dimension 1 and is equal to $A(z) + B(z)$. The tail vector $T(z)$ also has dimension 1 and is equal to 1. $M(z)$, the matrix of the body of the graph is the 1×1 matrix $M(z) = [A(z) + B(z)]$.

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Examples 8 and 9 show that the same class of combinatorial sequences may be associated with multiple transfer matrices. There are several ways to construct (and interpret) a weighted generating function.

Example 10. Let \mathcal{A} and \mathcal{B} be mutually exclusive classes of combinatorial objects with respective weighted generating function $A(z)$ and $B(z)$. Sequences of objects from \mathcal{A} or \mathcal{B} that end with \mathcal{B} but without two consecutive objects from \mathcal{B} can be thought of as walks on the graph below.



The edge with weighted generating function 1 from the head vertex to the tail vertex is no longer present because the empty object ε does not end with \mathcal{B} . The transfer matrix of this graph is the 4×4 matrix

$$M_*(z) = \begin{bmatrix} \circ & \mathcal{A} & \mathcal{B} & \bullet \\ \circ & 0 & A(z) & B(z) & 0 \\ \mathcal{A} & 0 & A(z) & B(z) & 0 \\ \mathcal{B} & 0 & A(z) & 0 & 1 \\ \bullet & 0 & 0 & 0 & 0 \end{bmatrix}.$$

The head vector $H(z)$ has dimension 2 and is equal to $(A(z), B(z))^\top$. The tail vector $T(z)$ also has dimension 2 and is equal to $(0, 1)^\top$. $M(z)$, the matrix of the body of the graph is the 2×2 matrix

$$\begin{bmatrix} \mathcal{A} & \mathcal{B} \\ \mathcal{A} & [A(z) & B(z)] \\ \mathcal{B} & [A(z) & 0] \end{bmatrix}.$$

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Let us now return to sequences of nucleotides without CG. To specify a transfer matrix, let us assume that G's and C's occur with frequency $p/2$ and that A's and T's occur with frequency $q/2$, where $q = 1 - p$. The weighted generating functions of the single nucleotides are thus $C(z) = G(z) = pz/2$ and $A(z) = T(z) = qz/2$. With these definitions, the transfer matrix of the graph shown in figure 2 is

$$\begin{aligned}
M_*(z) &= \begin{matrix} \circ & A & C & G & T & \bullet \\ \circ & 0 & A(z) & C(z) & G(z) & T(z) & 1 \\ A & 0 & A(z) & C(z) & G(z) & T(z) & 1 \\ C & 0 & A(z) & C(z) & 0 & T(z) & 1 \\ G & 0 & A(z) & C(z) & G(z) & T(z) & 1 \\ T & 0 & A(z) & C(z) & G(z) & T(z) & 1 \\ \bullet & 0 & 0 & 0 & 0 & 0 & 0 \end{matrix} \\
&= z/2 \begin{bmatrix} 0 & q & p & p & q & 1 \\ 0 & q & p & p & q & 1 \\ 0 & q & p & 0 & q & 1 \\ 0 & q & p & p & q & 1 \\ 0 & q & p & p & q & 1 \\ 0 & 0 & 0 & 0 & 0 & 0 \end{bmatrix}.
\end{aligned}$$

We still need to find the weighted generating function. Proposition 3 below shows how this is done.

Lemma 1. *Given a transfer graph and its transfer matrix $M_*(z)$, the weighted generating function of paths of n segments from vertex i to vertex j is the entry of $M_*(z)^n$ at position (i, j) .*

Proof. Proceed by induction. For $n = 1$, this is the definition of the transfer matrix. Assume that the property holds for some $n \geq 1$. A path of $n + 1$ segments from vertex i to vertex j is a path of n segments from vertex i to some vertex k , followed by a single segment from vertex k to vertex j . Using the induction hypothesis and taking the union on all vertices $1 \leq k \leq m + 2$, the weighted generating function of such paths is

$$\sum_{k=1}^{m+2} (M(z)^n)_{[i,k]} \times M(z)_{[k,j]} = (M(z)^{n+1})_{[i,j]},$$

which concludes the proof by induction. \square

Proposition 3. *The weighted generating function of the sequences generated by a transfer matrix $M_*(z)$ is the top right entry of the matrix $(I - M_*(z))^{-1}$, assuming that all the eigenvalues of $M(z)$ have modulus less than 1.*

Proof. From lemma 1, the weighted generating function is the top right entry of the matrix

$$I + M_*(z) + M_*(z)^2 + M_*(z)^3 + \dots$$

Observe that $(I - M_*(z)) \cdot (I + M_*(z) + M_*(z)^2 + \dots + M_*(z)^n) = I - M_*(z)^{n+1}$. If all the eigenvalues of $M(z)$ have modulus less than 1, the right hand side converges to the identity matrix I as n goes to infinity. This translates to

$$I + M_*(z) + M_*(z)^2 + M_*(z)^3 + \dots = (I - M_*(z))^{-1},$$

which concludes the proof. \square

Example 11. Let us continue example 7 where

$$M_*(z) = \begin{array}{c|ccc} & \circ & \mathcal{A} & \bullet \\ \circ & 0 & A(z) & 1 \\ \mathcal{A} & 0 & A(z) & 1 \\ \bullet & 0 & 0 & 0 \end{array}.$$

The weighted generating function of sequences of objects from \mathcal{A} is the top right entry of the matrix

$$(I - M_*(z))^{-1} = \frac{1}{1 - A(z)} \begin{bmatrix} 1 - A(z) & A(z) & 1 \\ 0 & 1 & 1 \\ 0 & 0 & 1 - A(z) \end{bmatrix}.$$

In other words, the weighted generating function of sequences of objects from \mathcal{A} is $1/(1 - A(z))$, consistently with proposition 2.

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Example 12. Let us continue examples 8 and 9. The first transfer matrix is

$$\begin{array}{c|cccc} & \circ & \mathcal{A} & \mathcal{B} & \bullet \\ \circ & 0 & A(z) & B(z) & 1 \\ \mathcal{A} & 0 & A(z) & B(z) & 1 \\ \mathcal{B} & 0 & A(z) & B(z) & 1 \\ \bullet & 0 & 0 & 0 & 0 \end{array}.$$

Applying proposition 3, the weighted generating function of sequences of objects from \mathcal{A} or \mathcal{B} is the top right entry of the matrix

$$\frac{1}{1 - A(z) - B(z)} \begin{bmatrix} 1 - A(z) - B(z) & A(z) & B(z) & 1 \\ 0 & 1 - B(z) & B(z) & 1 \\ 0 & A(z) & 1 - A(z) & 1 \\ 0 & 0 & 0 & 1 \end{bmatrix}.$$

The second transfer matrix is

$$\begin{array}{c|ccc} & \circ & \mathcal{A} \cup \mathcal{B} & \bullet \\ \circ & 0 & A(z) + B(z) & 1 \\ \mathcal{A} \cup \mathcal{B} & 0 & A(z) + B(z) & 1 \\ \bullet & 0 & 0 & 0 \end{array}.$$

Applying proposition 3 again, the weighted generating function of sequences of objects from from \mathcal{A} or \mathcal{B} is the top right entry of the matrix

$$\frac{1}{1 - A(z) - B(z)} \begin{bmatrix} 1 - A(z) - B(z) & A(z) + B(z) & 1 \\ 0 & 1 & 1 \\ 0 & 0 & 1 \end{bmatrix}.$$

Both terms are equal to $1/(1-A(z)-B(z))$. This illustrates that there are different ways to construct a sequence, some simpler than others.

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We now introduce a simplified way to find weighted generating functions from transfer matrices.

Proposition 4. *Given a transfer matrix*

$$M_*(z) = \begin{bmatrix} 0 & H(z)^\top & \psi(z) \\ 0 & M(z) & T(z) \\ 0 & 0 & 0 \end{bmatrix},$$

where $M(z)$ is a $m \times m$ matrix, $H(z)$ and $T(z)$ are vectors of dimension m and $\psi(z)$ has dimension 1, the weighted generating function of the sequences generated by $M_*(z)$ is

$$\psi(z) + H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z), \quad (3)$$

assuming that all the eigenvalues of $M(z)$ have modulus less than 1.

Proof. We need to compute the top-right term of the matrix $(I - M_*(z))^{-1}$. Using the matrix inversion formula with the cofactor matrix, this term is equal to $(-1)^{m+2}C/\det(I - M_*(z))$, where C is the determinant

$$\begin{vmatrix} -H(z)^\top & -\psi(z) \\ I - M(z) & -T(z) \end{vmatrix}.$$

Developing $\det(I - M_*(z))$ along the first column and then along the last row, we obtain $\det(I - M_*(z)) = (-1)^m \det(I - M(z))$. Developping C along the first row and then along the last column, we obtain

$$C = (-1)^m \psi(z) \det(I - M(z)) + \sum_{i=1}^m \sum_{j=1}^m H_i(z) (-1)^{i+j} C_{i,j}(z) T_j(z),$$

where $C_{i,j}$ is the cofactor of $I - M(z)$ at position (i, j) . Using once more the matrix inversion formula, we obtain

$$\frac{(-1)^{m+2}C}{\det(I - M_*(z))} = \psi(z) + H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z).$$

□

Example 13. Let us continue example 10 where

$$M_*(z) = \begin{array}{c|cccc} & \circ & \mathcal{A} & \mathcal{B} & \bullet \\ \circ & \begin{bmatrix} 0 & A(z) & B(z) & 0 \\ 0 & A(z) & B(z) & 0 \\ 0 & A(z) & 0 & 1 \\ 0 & 0 & 0 & 0 \end{bmatrix} \\ \mathcal{A} \\ \mathcal{B} \\ \bullet \end{array}.$$

We have seen that $H(z) = (A(z), B(z))^\top$, $T(z) = (0, 1)^\top$, $\psi(z) = 0$ and

$$M(z) = \begin{bmatrix} \mathcal{A} & \mathcal{B} \\ \mathcal{A} & \mathcal{B} \end{bmatrix} \begin{bmatrix} A(z) & B(z) \\ A(z) & 0 \end{bmatrix}.$$

Using proposition 4, the weighted generating function of the sequences generated by $M_*(z)$ is

$$0 + [A(z) \ B(z)] \cdot \frac{1}{1 - A(z)(1 + B(z))} \begin{bmatrix} 1 & B(z) \\ A(z) & 1 - A(z) \end{bmatrix} \cdot \begin{bmatrix} 0 \\ 1 \end{bmatrix} = \frac{B(z)}{1 - A(z)(1 + B(z))}.$$

*

We will sometimes represent transfer graphs as their bodies (associated with the matrix $M(z)$), and specify separately $\psi(z)$, $H(z)$ and $T(z)$. This will simplify the graphical representations, and using proposition 4 instead of proposition 3 will simplify the computations.

Now returning to sequences of nucleotides without CG, we have $\psi(z) = 1$, $H(z) = (A(z), C(z), G(z), T(z))^\top = (qz/2, pz/2, pz/2, qz/2)$, $T(z) = (1, 1, 1, 1)^\top$, and

$$M(z) = z/2 \begin{bmatrix} q & p & p & q \\ q & p & 0 & q \\ q & p & p & q \\ q & p & p & q \end{bmatrix}.$$

The inverse of $(I - M(z))^{-1}$ is

$$\frac{1}{\lambda(z)} \begin{bmatrix} (pz)^2 + 2qz + 4 & 2pz & pz(pz - 2) & 2qz \\ -pqz^2 + 2qz & -2(1 + q)z + 4 & 2pqz^2 & -pqz^2 + 2qz \\ 2qz & 2pz & -2(1 + q) + 4 & 2qz \\ 2qz & 2pz & pz(1 - pz) & (pz)^2 - 2(p + 1) + 4 \end{bmatrix}$$

where $\lambda(z) = 4 - 4z + (pz)^2$ is the determinant of $I - M(z)$. Now applying proposition 4, the weighted generating function of sequences of nucleotides without CG comes out as

$$1 + H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z) = \frac{4}{4 - 4z + (pz)^2}. \quad (4)$$

In this section we have seen that transfer matrices allow us to find the weighted generating functions of potentially intricate sequences of combinatorial objects. In the next section we will see how we can compute the probabilities of such sequences from their weighted generating function.

2.3 Asymptotic estimates

We know from expression (2) that we can recover the total weight (*i.e.* the probability) of objects of size k from the Taylor expansion of their weighted generating function. We will often need to extract a_k in expressions of the form $\sum_{k=1}^{\infty} a_k z^k$ so we introduce the following symbol.

Definition 4. Let $W(z)$ be a weighted generating function. The expression

$$[z^k]W(z)$$

stands for the **coefficient of z^k in W** and is equal to a_k , assuming that $W(z) = \sum_{k=1}^{\infty} a_k z^k$ on some neighborhood of 0.

The coefficient of z^k in a weighted generating function is usually a quantity of interest. For instance, in expression (4) it is the probability that a nucleotide sequence of size k does not contain any CG.

One of the main motivations for the analytic combinatorics approach is that we can efficiently approximate the coefficients of weighted generating functions. Proposition 5 below is a simplified version of the founding theorem of the field [4].

Proposition 5. If a function $W(z)$ is the ratio of two polynomials $P(z)/Q(z)$, and Q has only simple roots with distinct moduli, then

$$[z^k]W(z) \sim -\frac{P(z_1)}{Q'(z_1)} \frac{1}{z_1^{k+1}}, \quad (5)$$

where z_1 is the root of Q with smallest modulus.

Expression (5) is an asymptotic approximation: it becomes more accurate as k increases, *i.e.* as we consider objects of increasing size. This is the most fundamental property of the general analytic combinatorics strategy.

The roots of Q are called “singularities” of the function W . They are values where the function is not defined. The roots of Q with multiplicity 1 (*i.e.* the values for which Q vanishes but its derivative does not) are also referred to as “simple poles” of W .

Proposition 5 says that the asymptotic growth of the coefficients of the series expansion of $W(z)$ is dictated by the singularity of smallest *modulus*, also known as the “dominant singularity”. We first state a lemma that will be useful to demonstrate proposition 5.

Lemma 2. If $|z| < a$, then

$$\frac{1}{1 - z/a} = \sum_{k=0}^{\infty} \frac{z^k}{a^k}. \quad (6)$$

Proof. Proceed as in example 5, replacing p by $1/a$. \square

We now prove proposition 5.

Proof. Let z_1, z_2, \dots, z_n be the complex roots of Q , sorted by increasing order of *modulus*. Since Q has only simple roots, there exists constants β_1, \dots, β_n such that the partial fraction decomposition of the rational function $P(z)/Q(z)$ can be written as

$$W(z) = P(z)/Q(z) = R(z) + \sum_{j=1}^n \frac{\beta_j}{z - z_j} = R(z) - \sum_{j=1}^n \frac{\beta_j/z_j}{1 - z/z_j}. \quad (7)$$

In the expression above, R is a polynomial that is nonzero if and only if the degree of P is higher than the degree of Q . Either way, the coefficient of z^k in $R(z)$ is 0 for k higher than the degree of R , so the coefficients of R do not contribute to $[z^k]W(z)$ for large k . We can thus assume $R(z) = 0$ without loss of generativity. From lemma 2 we can expand each of the n terms of the sum as

$$-\frac{\beta_j/z_j}{1 - z/z_j} = -\beta_j/z_j \sum_{k=0}^{\infty} \frac{z^k}{z_j^k}.$$

Substituting the expression above in (7), we obtain

$$[z^k]W(z) = -\sum_{j=1}^n \frac{\beta_j}{z_j^{k+1}}. \quad (8)$$

Since z_1 is the root with smallest *modulus*, the sum above is dominated by the term $1/z_1^{k+1}$ as k increases, so the coefficient of z^k in $W(z)$ is asymptotically equivalent to

$$[z^k]W(z) \sim -\frac{\beta_1}{z_1^{k+1}}.$$

To find the value of β_1 , we factorize $Q(z)$ as $(z - z_1)Q_1(z)$, which is possible because z_1 is a root of Q , and we write

$$\frac{P(z)}{Q(z)} = \frac{P(z)}{(z - z_1)Q_1(z)} = \frac{\beta_1}{z - z_1} + \sum_{j=2}^n \frac{\beta_j}{z - z_j}.$$

Multiplying through by $(z - z_1)$ and setting $z = z_1$, we obtain the expression $\beta_1 = P(z_1)/Q_1(z_1)$. Differentiating the expression $Q(z) = (z - z_1)Q_1(z)$ shows that $Q'(z_1) = Q_1(z_1)$, and thus that $\beta_1 = P(z_1)/Q'(z_1)$, which concludes the proof. \square

Remark 7. Expression (8) is not an approximation, it is the exact value of the coefficient of z^k in $W(z) = P(z)/Q(z)$. By keeping more than one term in the sum, we can obtain more accurate estimates, and by keeping all the terms we obtain the exact value.

Remark 8. The asymptotic estimate (5) converges exponentially fast to the coefficient as k increases. To see this, divide the exact expression (8) by its leading term $-\beta_1/z_1^{k+1}$ and obtain

$$\frac{-\sum_{j=1}^n \beta_j/z_j^{k+1}}{-\beta_1/z_1^{k+1}} = 1 + \sum_{j=2}^n \frac{\beta_j}{\beta_1} \left(\frac{z_1}{z_j} \right)^{k+1}.$$

Since $|z_1| < |z_j|$ for $2 \leq j \leq n$, the error terms are $O(|z_1/z_2|^k)$ so they decrease exponentially fast as k increases.

Remark 9. Proposition 5 does not hold if z_1 is not a simple pole. The proof can be generalized, but the resulting asymptotic formula is different. Proposition 10 in section 5.2 shows the coefficient asymptotics for poles of second order.

Recall from section 2.2 that the weighted generating function of nucleotide sequences without CG is

$$W(z) = \frac{4}{4 - 4z + (pz)^2}. \quad (4)$$

Here $Q(z) = 4 - 4z + (pz)^2$ has two distinct roots, $z_1 = 2(1 - \sqrt{1 - p^2})/p^2$ and $z_2 = 2(1 + \sqrt{1 - p^2})/p^2$. Since $Q'(z) = 2p^2z - 4$, proposition 5 yields

$$\begin{aligned} [z^k]W(z) &\sim -\frac{P(z_1)}{Q'(z_1)} \frac{1}{z_1^{k+1}} = \frac{1}{\sqrt{1-p^2}} \left(\frac{p^2/2}{1 - \sqrt{1-p^2}} \right)^{k+1} \\ &= \frac{1}{\sqrt{1-p^2}} \left(\frac{1 + \sqrt{1-p^2}}{2} \right)^{k+1}. \end{aligned}$$

In this case we can also obtain the exact value by using the second singularity. The probability that a sequence of size k contains no CG is exactly equal to

$$\begin{aligned} &-\frac{P(z_1)}{Q'(z_1)} \frac{1}{z_1^{k+1}} - \frac{P(z_2)}{Q'(z_2)} \frac{1}{z_2^{k+1}} \\ &= \frac{1}{\sqrt{1-p^2}} \left[\left(\frac{p^2/2}{1 - \sqrt{1-p^2}} \right)^{k+1} - \left(\frac{p^2/2}{1 + \sqrt{1-p^2}} \right)^{k+1} \right] \\ &= \frac{1}{\sqrt{1-p^2}} \left[\left(\frac{1 + \sqrt{1-p^2}}{2} \right)^{k+1} - \left(\frac{1 - \sqrt{1-p^2}}{2} \right)^{k+1} \right]. \end{aligned}$$

If all the nucleotides have the same frequency, then $p = 1/2$ and the probability is approximately equal to $(0.9330127^{k+1} - 0.0669873^{k+1})/1.154701$. From $k = 5$, the second term of the sum is over one million times smaller than the first term.

This example illustrates the elegance and the efficiency of the analytic combinatorics approach. It also shows how accurate the approximate solution can be.

In conclusion, we summarize the analytic combinatorics strategy as follows: (*i*) define simple objects associated with simple generating functions, (*ii*) combine these objects into more complex structures, (*iii*) translate those combinations into more complex generating functions, and (*iv*) extract coefficients using approximation theorems such as use proposition 5.

3 Exact seeding

3.1 Reads, error symbols and error-free intervals

From the experimental point of view, a sequencing read is the result of an assay on some polymer of nucleic acid. The output of the assay is the decoded sequence of monomers that compose the molecule. Three types of sequencing errors can occur: *substitutions*, *deletions* and *insertions*. A substitution is a nucleotide that is different in the molecule and in the read, a deletion is a nucleotide that is present in the molecule but not in the read, and an insertion is a nucleotide that is absent in the molecule but present in the read.

For our purpose, the focus is not the nucleotide sequence *per se*, but whether the nucleotides correct. Thus, we need only four symbols to describe a read: one for each type or error, plus one for correct nucleotides. In this view, a read is a finite sequence of letters from an alphabet of four symbols.

Errors in the read are initially unknown. In practice, the only way to detect them is to align the read with the reference sequence in order to highlight the differences. This operation may be difficult to perform, especially if the read has many errors. But here, we are only concerned with computing the probabilities of occurrence of certain reads, so we will assume that the errors are known.

A read can be seen as a sequence of symbols. Figure 3 shows the typical structure of a read, together with the symbols that we will use for correct nucleotides, substitutions, deletions, and insertions.



Figure 3: **Read as sequence of error symbols.** Reads consist of correct nucleotides (white boxes), substitutions (black boxes), deletions (vertical bars) and insertions (grey boxes).

A read can be partitioned uniquely into maximal sequences of identical symbols called “intervals”. Intervals of correct nucleotides will have a particular importance, so we introduce the somewhat simpler term “error-free interval”.

Definition 5. An *error-free interval* is a sequence of correct nucleotides that cannot be extended left or right.

Instead of sequences of symbols, reads can be seen as sequences of either error-free intervals or error symbols (see figure 4). We will use this construction throughout section 3. It will allow us to control the size of the largest error-free interval (see below).



Figure 4: **Read as sequence of error-free intervals or error symbols.** Consecutive correct nucleotides can be lumped together in error-free intervals. There is a unique representation of a read as a sequence of either error-free intervals or error symbols. The error-free intervals are highlighted in this example, together with their size.

These formal definitions will allow us to specify the weighted generating functions of the reads of interest and then approximate their probability of occurrence using the results of section 2.

3.2 Exact seeds

After sequencing, one of the most common operations is to map the reads in a reference genome, *i.e.* find the identity of the fragment that was sequenced from a list of candidates. The exact search problem can be solved efficiently, provided the right indexing data structures are available, but sequencing errors make the task more difficult.

In most mapping algorithms, the search starts by looking for perfect matches between a fragment of the read and a sequence from the genome. This match is called a “seed”, and it allows to quickly eliminate most of the search space in order to focus on a few promising candidate sequences.

This strategy greatly accelerates the mapping process, but it is not failsafe. Seeds are useful only if they are error-free; otherwise they will either match nothing, or match the wrong sequence. This means that error-free intervals hold the key to the problem. If there exists a long enough error-free interval, it may be used as a seed to find the identity of the read. Here we will use a definition of “seed” that corresponds to this intuition.

Definition 6. An *exact γ -seed* is an error-free interval of size at least γ .

Remark 10. In what follows, we will often refer to an “exact γ -seed” as a “seed” when the concrete value of γ is either clear from the context or irrelevant.

The main goal of section 3 is to develop models to estimate the probability that a read contains an exact γ -seed. Our strategy is to construct the weighted generating functions of reads that *do not* contain any exact γ -seed. For this, we will decompose such reads as sequences of either error symbols or error-free intervals of size less than γ . We will obtain their weighted generating functions

from proposition 4 and use proposition 5 to approximate their probability of occurrence.

We will find the weighted generating function of all the reads $R(z)$, and then the weighted generating function of reads without exact γ -seeds $S_\gamma(z)$. The probability that a read of size k has no exact γ -seed will then be equal to the total weight of reads of size k without seed, divided by the total weight of reads of size k , *i.e.*

$$\frac{[z^k]S_\gamma(z)}{[z^k]R(z)}. \quad (9)$$

To introduce the concepts progressively, we will first describe simplified models where some types of errors are disallowed, and gradually increase the complexity towards the model with all error types.

3.3 Substitutions only

In the simplest model, we assume that errors can be only substitutions, and that they occur with the same probability p for every nucleotide. We will refer to this model as the “uniform substitutions error model”. Due to its simplicity, we will be able to go at greater depth and obtain more advanced results with this model (especially in the later sections). Importantly, the model is not overly simple and it has some real practical applications. For instance, it describes reasonably well the error model of the Illumina platforms (where p is around 0.01).

We will first use proposition 4 to obtain the weighted generating function $R(z)$ of all reads under this error model, from which we are going to deduce the weighted generating function $S_\gamma(z)$ of reads without exact γ -seeds. We will then use proposition 5 to derive an approximation formula for the probability that a read does not contain exact γ -seeds.

Under the uniform substitutions model, reads are sequences of either substitutions or error-free intervals. They can be thought of as a walk on the graph shown in figure 5. The symbol Δ_0 stands for error-free intervals and the symbol S stands for single substitutions. $F(z)$ and pz indicate the weighted generating functions of error-free intervals and substitutions, respectively. The fact that an error-free interval cannot follow another error-free interval is a consequence of the definition: two consecutive intervals must be merged into a larger interval.

Remark 11. *In all the error models described in section 3, the error-free intervals cannot have size 0. The reason is that a read must correspond to exactly one path in the transfer graph and that sequences of combinatorial classes containing the empty object ε are not uniquely defined (see remark 4).*

A substitution is a single nucleotide and thus has size 1. Because substitutions have probability p , their weighted generating function is pz . Conversely, the weighted generating function of correct nucleotides is qz . Error-free intervals are non-empty sequences of correct nucleotides, so by proposition 2 their weighted generating function is

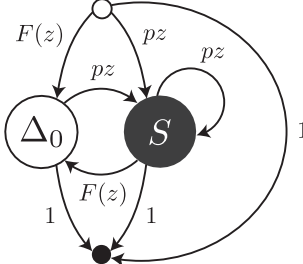


Figure 5: **Transfer graph of reads with uniform substitutions.** Reads are viewed as sequences of error-free intervals (symbol Δ_0) or substitutions (symbol S). $F(z)$ and pz are the weighted generating functions of error-free intervals and individual substitutions, respectively.

$$F(z) = qz + (qz)^2 + (qz)^3 + \dots = \frac{qz}{1 - qz}. \quad (10)$$

The transfer matrix associated with the body of the transfer graph shown in figure 5 is

$$M(z) = \begin{bmatrix} \Delta_0 & S \\ S & F(z) \end{bmatrix} = \begin{bmatrix} 0 & pz \\ F(z) & pz \end{bmatrix}.$$

A read can start with an error-free interval or with a substitution, so the head vector $H(z)$ is equal to $(F(z), pz)^\top$. Similarly, a read can end with an error free interval or with a substitution, so the tail vector $T(z)$ is equal to $(1, 1)^\top$. Here $\psi(z) = 1$, as the only read excluded by our definition is the empty sequence ε . Applying proposition 4, the weighted generating function of reads is $R(z) = \psi(z) + (F(z), pz)^\top \cdot (I - M(z))^{-1} \cdot T(z)$, or

$$R(z) = 1 + [F(z) \ pz] \cdot \frac{1}{\lambda(z)} \begin{bmatrix} 1 - pz & pz \\ F(z) & 1 \end{bmatrix} \cdot \begin{bmatrix} 1 \\ 1 \end{bmatrix},$$

where $\lambda(z) = 1 - pz(1 + F(z))$ is the determinant of $I - M(z)$. Finishing the computations, we obtain

$$R(z) = \frac{1 + F(z)}{1 - pz(1 + F(z))} = \frac{1}{1 - z}. \quad (11)$$

Since $1/(1 - z) = 1 + z + z^2 + \dots$, this means that for any $k \geq 0$, the total weight of reads of size k is equal to 1. As a consequence, ratio (9) reduces to $[z^k]S_\gamma(z)$.

To find the weighted generating function of reads without any exact γ -seeds, we need to limit error-free intervals to a maximum size of $\gamma - 1$. To do this, we

can replace $F(z)$ in (11) by its truncation $F_\gamma(z) = qz + (qz)^2 + \dots + (qz)^{\gamma-1}$. We obtain

$$S_\gamma(z) = \frac{1 + F_\gamma(z)}{1 - pz(1 + F_\gamma(z))} = \frac{1 + qz + \dots + (qz)^{\gamma-1}}{1 - pz(1 + qz + \dots + (qz)^{\gamma-1})}. \quad (12)$$

Remark 12. Viewing reads as sequences of either error-free intervals or substitutions was important to obtain the weighted generating function of reads without seeds. It allowed us to just replace $F(z)$ by $F_\gamma(z)$ to find the expression immediately.

The task is now to extract the coefficient of z^k in $S_\gamma(z)$. As explained in section 2.3, we look for an asymptotic estimate using proposition 5. For this, we need to find the singularities of $S_\gamma(z)$.

It is interesting to look in detail into the problem of finding the singularities of $S_\gamma(z)$. To highlight some general features of the problem, we start with a concrete case. The left panel of Figure 6 shows the values of the denominator of $S_\gamma(z)$ with $p = 0.1$ and $\gamma = 17$. S_{17} has one real root greater than 1. The remaining singularities of S_{17} are complex and they seem to be evenly spaced on the same circle, as can be seen on the right panel of Figure 6. This is only a visual impression. In fact the singularities are not exactly on the same circle and their rotation angles are not exactly regular.

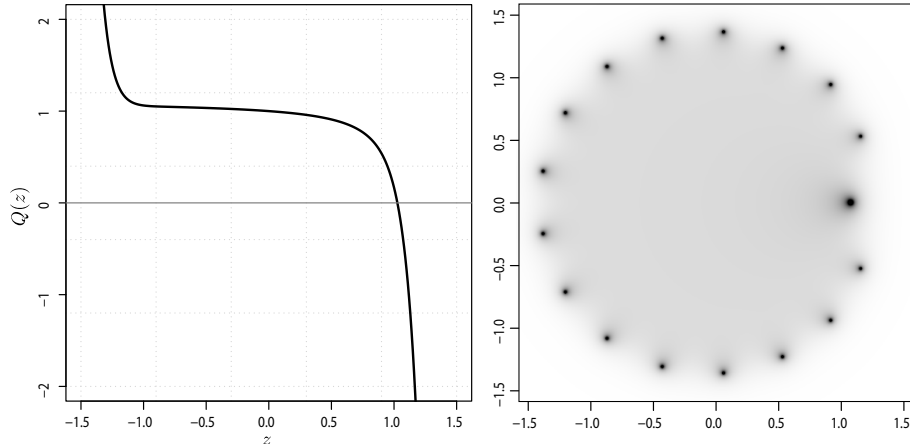


Figure 6: **Singularities of S_{17} .** $Q(z)$ denotes the denominator of $S_{17}(z)$ from expression (12) with $p = 0.1$ and $\gamma = 17$. *Left:* The value of $Q(z)$ is represented for z real by the bold line. *Right:* The value of $|1/Q(z)|$ is represented for z complex by the heat map on the complex plane. Darker pixels correspond to higher values. Sixteen singularities of S_{17} lie close to a circle. The remaining seventeenth is the one shown on the left panel. It is the dominant singularity because it is the closest to the origin.

It is “fortunate” that the dominant root of S_{17} is a real number because we can use efficient numerical methods to approximate it (*e.g.* bisection or the Newton-Raphson method). The following proposition shows that this is no

accident: the dominant singularity of S_γ is always a real positive number greater than 1.

Proposition 6. *S_γ as expressed in (12) has exactly one positive real singularity. This is the dominant singularity and it is greater than 1.*

Proof. Write $S_\gamma(z) = P(z)/Q(z)$ and search the roots of the denominator $Q(z) = 1 - pz(1 + qz + \dots + (qz)^{\gamma-1})$. First, we show that Q has exactly one positive real root greater than 1. For real $z > 0$, pz and $1 + qz + \dots + (qz)^{\gamma-1}$ are strictly increasing, so $Q(z)$ is strictly decreasing. Since $Q(1) = q^d > 0$ there is no root in the interval $(0, 1)$ and since $\lim_{z \rightarrow \infty} Q(z) = -\infty$, Q vanishes for a real number greater than 1.

Second, we show that this is the root with smallest *modulus*. Express the complex roots of Q as $Re^{i\theta}$. They satisfy the equation

$$1 - pRe^{i\theta} \frac{1 - q^\gamma R^\gamma e^{i\gamma\theta}}{1 - qRe^{i\theta}} = 0, Re^{i\theta} \neq \frac{1}{q}.$$

Multiplying through by $1 - qRe^{i\theta}$, we obtain an equation of which we separate the real and the imaginary parts to obtain

$$\begin{cases} pR \cos(\theta) - 1 = pq^\gamma R^{\gamma+1} \cos((\gamma+1)\theta) \\ pR \sin(\theta) = pq^\gamma R^{\gamma+1} \sin((\gamma+1)\theta) \end{cases} \quad Re^{i\theta} \neq \frac{1}{q}.$$

Squaring and summing, we obtain the following equation

$$p^2 R^2 = (pq^\gamma R^{\gamma+1})^2 + 2pR \cos(\theta) - 1. \quad (13)$$

Considering this an equation of $R > 0$, the solution is minimal when $2pR \cos(\theta)$ is maximal, *i.e.* when $\theta = 0$. In other words, if there is a positive real root, its *modulus* is the minimum among the roots. We have seen above that there exists exactly one, so it is the dominant singularity of S_γ . \square

We now have all the tools to approximate the coefficients of $S_\gamma(z)$ using proposition 5 and obtain the approximate probability that a read contains no seed.

Proposition 7. *The probability that a read of size k has no seed under the uniform substitutions model is asymptotically equivalent to*

$$\frac{C}{z_1^{k+1}},$$

where z_1 is the only real positive root of $1 - pz(1 + qz + \dots + (qz)^{\gamma-1})$, and where

$$C = \frac{(1 - qz_1)^\gamma}{p^2 z_1 (1 - (\gamma+1 - \gamma q z_1)(q z_1)^\gamma)}. \quad (14)$$

Proof. Apply propositions 5 to $S_\gamma(z)$ and then proposition 6, together with the fact that for any singularity z of S_γ we have $1 + qz + \dots + (qz)^{\gamma-1} = 1/pz$. \square

We now illustrate proposition 7 with a concrete example explaining how the calculations are done in practice.

Example 14. Let us approximate the probability that a read of size $k = 100$ has no seed for $\gamma = 17$ and for a substitution rate $p = 0.1$. To find the dominant singularity of S_{17} , we need to solve $1 - 0.1z \times (1 + 0.9z + \dots + (0.9z)^{16}) = 0$. We rewrite the equation as $1 - 0.1z \times (1 - (0.9z)^{17})/(1 - 0.9z) = 0$ and use bisection to solve it numerically, yielding $z_1 \approx 1.0268856$. Substituting this value in (14) yields $C \approx 1.396145$, so the probability that a read contains no seed is approximately $1.396145/1.0268856^{101} \approx 0.095763$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.09575 - 0.09577$. The computational cost of the analytic combinatorics approach is infinitesimal compared to the random simulations, and the precision is much higher for $k = 100$.

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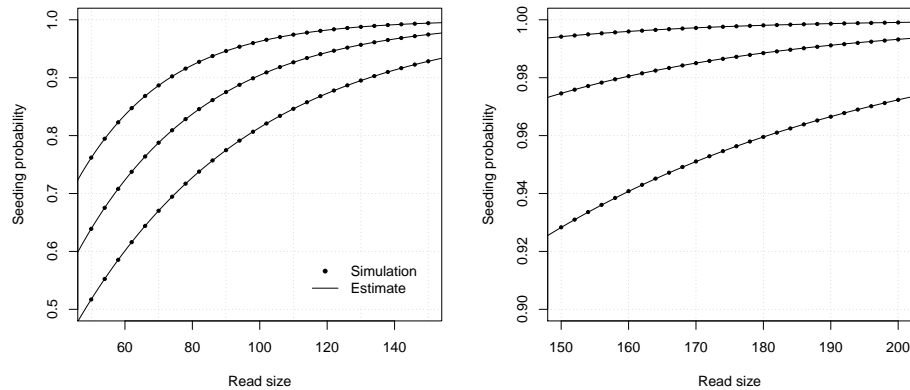


Figure 7: **Example estimates for substitutions only.** The analytic combinatorics estimates of proposition 7 are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size contains a seed, either estimated by 10,000,000 random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$ and $p = 0.08$, $p = 0.10$ or $p = 0.12$ (from top to bottom).

Overall, the analytic combinatorics estimates are close to the exact values. Figure 7 illustrates the precision of the estimates for different values of the error rate p and of the read size k .

3.4 Substitutions and deletions

To not jump too fast into the difficulties, we will now describe a semi-realistic model where errors can be deletions or substitutions, but not insertions. As in the case of uniform substitutions, we assume that every nucleotide call is false

with a probability p and true with a probability $1 - p = q$. Here, we also assume that the “space” between consecutive nucleotides can contain a deletion with probability δ .

A deletion may be adjacent to a substitution, or lie in between two correct nucleotides. In the first case, the deletion does not interrupt any error-free interval so it does not change the probability that the read contains a seed. For this reason, we ignore deletions next to substitutions. More precisely, we assume that they can occur, but whether they do has no importance for the problem.

Under this error model, a read can be thought of as a walk on the graph shown in figure 8. The graph is almost the same as the one shown figure 5; the only difference is the edge labelled $\delta F(z)$ from Δ_0 to Δ_0 . This edge represents the fact that under this error model, an error-free interval can follow another one if a deletion with weighted generating function δ is present in between (as illustrated for instance in figure 4).

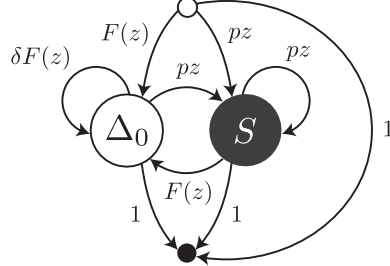


Figure 8: Transfer graph of reads with uniform substitutions and deletions. Reads are viewed as sequences of error-free intervals (symbol Δ_0) or substitutions (symbol S). An error-free interval can follow another one if a deletion is present in between. $F(z)$ and pz are the weighted generating functions of error-free intervals and individual substitutions, respectively. $\delta F(z)$ is the weighted generating function of a deletion followed by an error-free interval.

The weighted generating function of error-free intervals $F(z)$ has a different expression from that of section 3.3. When the size of an error-free interval is 1, the weighted generating function is just qz . For a size $k > 1$, there are $k - 1$ “spaces” between the nucleotides, so the weighted generating function is $(1 - \delta)^{k-1}(qz)^k$. Summing for all the possible sizes, we obtain the weighted generating function of error-free intervals as

$$F(z) = qz + (1 - \delta)(qz)^2 + (1 - \delta)^2(qz)^3 + \dots = \frac{qz}{1 - (1 - \delta)qz}. \quad (15)$$

The transfer matrix associated with the body of the transfer graph shown in figure 8 is

$$M(z) = \begin{matrix} & \Delta_0 & S \\ \Delta_0 & \begin{bmatrix} \delta F(z) & pz \\ F(z) & pz \end{bmatrix} \\ S & \end{matrix}.$$

The head and tail vectors are the same as in section 3.3, *i.e.* $H(z) = (F(z), pz)^\top$, $T(z) = (1, 1)^\top$ and $\psi(z) = 1$. Applying proposition 4 yields

$$R(z) = (I - M(z))^{-1} = 1 + [F(z) \ pz] \cdot \frac{1}{\lambda(z)} \begin{bmatrix} 1 - pz & pz \\ F(z) & 1 - \delta F(z) \end{bmatrix} \cdot \begin{bmatrix} 1 \\ 1 \end{bmatrix},$$

where $\lambda(z) = 1 - pz - (pz(1 - \delta) + \delta)F(z)$ is the determinant of $I - M(z)$. Finishing the computations we obtain

$$R(z) = \frac{1 + (1 - \delta)F(z)}{1 - pz - (pz(1 - \delta) + \delta)F(z)} = \frac{1}{1 - z}. \quad (16)$$

As in section 3.3 the result is $1/(1 - z) = 1 + z + z^2 + \dots$, which means that the total weight of reads of size k is equal to 1 for every $k \geq 0$. This means that ratio (9) reduces one again to $[z^k]S_\gamma(z)$.

To find the weighted generating function of reads without any exact γ -seeds, we need to bound the size of error-free intervals to a maximum of $\gamma - 1$, *i.e.* to replace $F(z)$ by its truncation $F_\gamma(z) = qz + (1 - \delta)(qz)^2 + \dots + (1 - \delta)^{\gamma-2}(qz)^{\gamma-1}$. With this definition, the weighted generating function of seedless reads is

$$S_\gamma(z) = \frac{1 + (1 - \delta)F_\gamma(z)}{1 - pz - (pz(1 - \delta) + \delta)F_\gamma(z)}. \quad (17)$$

Applying proposition 5 to this expression, we obtain the following approximation formula.

Proposition 8. *The probability that a read of size k has no seed under the model of uniform substitutions and deletions is asymptotically equivalent to*

$$\frac{C}{z_1^{k+1}},$$

where z_1 is the only real positive root of the denominator of $S_\gamma(z)$, *i.e.* the root of $1 - pz - (pz(1 - \delta) + \delta)(qz + (1 - \delta)(qz)^2 + \dots + (1 - \delta)^{\gamma-2}(qz)^{\gamma-1})$, and

$$C = \frac{z_1(1 - (1 - \delta)(1 - p)z_1)^2}{((p + q\delta)z_1 - c_1(1 - \delta)^{\gamma-1}(qz_1)^\gamma)(\delta + (1 - \delta)pz_1)}, \text{ where} \quad (18)$$

$$c_1 = \gamma\delta - (1 - \delta)((\gamma - 1)\delta - p((\gamma - 1)\delta + \gamma + 1))z_1 - \gamma(1 - \delta)^2pqz_1^2.$$

We illustrate proposition 8 with a concrete example explaining how the calculations are done in practice.

Example 15. Let us approximate the probability that a read of size $k = 100$ has no seed for $\gamma = 17$, $p = 0.05$ and $\delta = 0.15$. To find the dominant singularity of S_{17} we solve $1 - 0.05z - (0.0425z + 0.15)(0.95z + 0.85(0.95z)^2 + \dots + 0.85^{15}(0.95z)^{16}) = 0$. We write it as $1 - 0.05z - (0.0425z + 0.15)(0.95z - 0.85^{16}(0.95z)^{17})/(1 - 0.8075z) = 0$ and use bisection, yielding $z_1 \approx 1.006705$. Now substituting the obtained value in (18) gives $C \approx 1.096177$, so the probability that a read contains no seed is approximately $1.096177/1.006705^{101} \approx 0.558141$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.55813 - 0.55816$.

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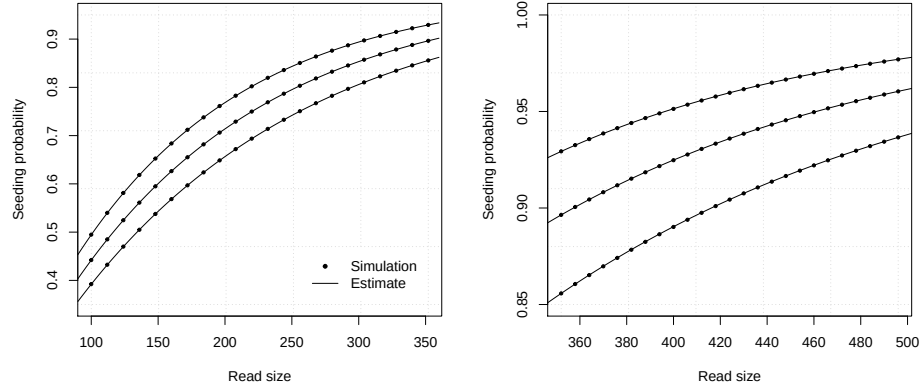


Figure 9: **Example estimates for substitutions and deletions.** The analytic combinatorics estimates of proposition 8 are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size contains a seed, either estimated by 10,000,000 random simulations (dots), or by proposition 8 (lines). The curves are drawn for $\gamma = 17$, $p = 0.05$ and $\delta = 0.14$, $\delta = 0.15$ or $\delta = 0.16$ (from top to bottom).

Once again, the analytic combinatorics estimates are close to the exact values. Figure 9 illustrates the precision of the estimates for different values of the deletion rate δ and of the read size k .

3.5 Substitutions, deletions and insertions

Here we consider the model that we will refer to as “full error model” because all types of errors are allowed. Introducing insertions brings two additional difficulties. The first is that a substitution is indistinguishable from an insertion followed by a deletion (or a deletion followed by an insertion). By convention, we will count all these cases as substitutions. As a consequence, a deletion can never be found next to an insertion. The second difficulty is that insertions

usually come in bursts. This is also the case of deletions, but we could neglect it because this does not affect the size of the interval (all deletions have size 0).

To model insertion bursts, we need to assign a probability r to the first insertion, and a probability $\tilde{r} > r$ to all subsequent insertions of the burst. We will still denote the probability of a substitution p and that of a correct call q , but here $p + q + r = 1$. We will also assume that an insertion burst stops with probability $1 - \tilde{r}$ at each position of the burst.

Under this error model, reads can be thought of as a walk on the graph shown in figure 10. The symbols Δ_0 , S and I stand for error-free intervals, single substitutions and single insertions, respectively. The body of the graph is represented separately from the head and tail vertices to avoid overloading the figure.

The terms $F(z)$, pz and $\delta F(z)$ are the same as in section 3.4. The terms rz and $\tilde{r}z$ are the weighted generating functions of the first inserted nucleotide and of all subsequent nucleotides of the insertion burst, respectively. The burst terminates with probability $1 - \tilde{r}$ and is followed by an error-free interval or by a substitution. The total weight of these two cases is $p + q < 1$, so we need to further scale the weighted generating functions by a factor $p + q = 1 - r$.

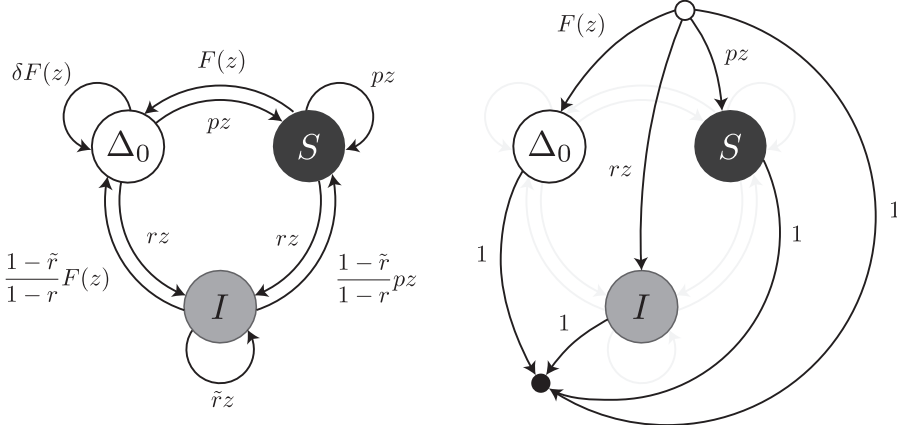


Figure 10: **Transfer graph of reads with full error model.** Reads are viewed as sequences of error-free intervals (symbol Δ_0), substitutions (symbol S) or insertions (symbol I). To not overload the figure, the body of the transfer graph is shown on the left, and the head and tail edges are shown on the right. $F(z)$ and pz are the weighted generating functions of error-free intervals and individual substitutions, respectively. $\delta F(z)$ is the weighted generating function of a deletion followed by an error-free interval. rz and $\tilde{r}z$ are the weighted generating functions of the first and all subsequent insertions of a burst, respectively. $(1 - \tilde{r})F(z)/(1 - r)$ is the weighted generating function of an error-free interval following an insertion and $(1 - \tilde{r})pz/(1 - r)$ is the weighted generating function of a substitution following an insertion.

The expression of the weighted generating function of error-free intervals

$F(z)$ is the same as in section 3.4, namely

$$F(z) = qz + (1 - \delta)(qz)^2 + (1 - \delta)^2(qz)^3 + \dots = \frac{qz}{1 - (1 - \delta)qz}.$$

The transfer matrix associated with the body of the transfer graph shown in figure 10 is

$$M(z) = \begin{matrix} & \Delta_0 & S & I \\ \Delta_0 & \delta F(z) & pz & rz \\ S & F(z) & pz & rz \\ I & \frac{1-\tilde{r}}{1-r}F(z) & \frac{1-\tilde{r}}{1-r}pz & \tilde{r}z \end{matrix}.$$

Here the head vector $H(z)$ is equal to $(F(z), pz, rz)^\top$, the tail vector $T(z)$ is equal to $(1, 1, 1)^\top$, and $\psi(z) = 1$. The expression of $(I - M(z))^{-1}$ is omitted because it is cumbersome, but according to proposition 4 all we need is the value of $\psi(z) + H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$, which is equal to

$$R(z) = \frac{(1 - r)(1 - (\tilde{r} - r)z)(1 + (1 - \delta)F(z))}{1 - a(z) - b(z)F(z)} = \frac{1}{1 - z}, \quad (19)$$

where $a(z)$ and $b(z)$ are second degree polynomials defined as

$$\begin{aligned} a(z) &= r + (1 - r)(p + \tilde{r})z - p(\tilde{r} - r)z^2, \text{ and} \\ b(z) &= \delta(1 - r) + ((1 - r)(p - \delta(p + \tilde{r}) + (1 - \tilde{r})r))z - p(1 - \delta)(\tilde{r} - r)z^2. \end{aligned} \quad (20)$$

Again, the terms cancel out and we obtain the simple expression $1/(1 - z) = 1 + z + z^2 + \dots$ where all the coefficients are equal to 1. This means that once again, ratio (9) reduces to $[z^k]S_\gamma(z)$. To find the weighted generating function of reads without exact γ -seed, we replace $F(z)$ in expression (19) by its truncated version

$$F_\gamma(z) = qz + (1 - \delta)(qz)^2 + (1 - \delta)^2(qz)^3 + \dots + (1 - \delta)^{\gamma-2}(qz)^{\gamma-1}.$$

We obtain the following expression

$$S_\gamma(z) = \frac{(1 - r)(1 - (\tilde{r} - r)z)(1 + (1 - \delta)F_\gamma(z))}{1 - a(z) - b(z)F_\gamma(z)}, \quad (21)$$

where $a(z)$ and $b(z)$ are defined as in (20).

Remark 13. Note that when $r = \tilde{r} = 0$, $a(z) = pz$ and $b(z) = pz(1 - \delta) + \delta$, so expression (21) becomes

$$S_\gamma(z) = \frac{1 + (1 - \delta)F_\gamma(z)}{1 - pz - (pz(1 - \delta) + \delta)F_\gamma(z)}.$$

This is expression (17), i.e. the model described in section 3.4. When we also have $\delta = 0$, this expression further simplifies to

$$S_\gamma(z) = \frac{1 + F_\gamma(z)}{1 - pz(1 + F_\gamma(z))}.$$

This is expression (12), i.e. the model described in section 3.3. In other words, the error models described previously are special cases of the full error model.

As in the previous sections, we can use proposition 5 to obtain asymptotic approximations for the probability that the reads contain no seed.

Proposition 9. *The probability that a read of size k has no seed under the full error model is asymptotically equivalent to*

$$\frac{C}{z_1^{k+1}},$$

where z_1 is the only real positive root of the polynomial $1 - a(z) - b(z)F_\gamma(z) = 1 - a(z) - b(z)(qz + (1 - \delta)(qz)^2 + \dots + (1 - \delta)^{\gamma-2}(qz)^{\gamma-1})$, and $C = c_1/c_2$, with

$$c_1 = (1 - (\tilde{r} - r)z_1) \frac{1 - ((1 - \delta)(qz))^\gamma}{1 - (1 - \delta)qz}$$

and

$$\begin{aligned} c_2 &= a'(z_1) - \left(b'(z_1) + \frac{(1 - \delta)q}{1 - (1 - \delta)qz} \right) \frac{qz - (1 - \delta)^{\gamma-1}(qz)^\gamma}{1 - (1 - \delta)qz} \\ &\quad + b(z_1) \frac{q - \gamma(1 - \delta)^{\gamma-1}q^\gamma z^{\gamma-1}}{1 - (1 - \delta)qz}. \end{aligned}$$

We illustrate proposition 9 with a concrete example explaining how the calculations are done in practice.

Example 16. Let us approximate the probability that a read of size $k = 100$ has no seed for $\gamma = 17$, $p = 0.05$, $\delta = 0.15$, $r = 0.05$ and $\tilde{r} = 0.45$. With these values, $a(z) = 0.05 + 0.475z - 0.02z^2$ and $b(z) = 0.1425 + 0.00375z - 0.017z^2$. We need to solve $0.95 - 0.475z + 0.02z^2 - (0.1425 + 0.00375z - 0.017z^2)(0.9z + 0.85(0.9z)^2 + \dots + 0.85^{15}(0.9z)^{16}) = 0$. We rewrite the equation as $0.95 - 0.475z + 0.02z^2 - (0.1425 + 0.00375z - 0.017z^2)(0.9z - 0.85^{15}(0.9z)^{16})/(1 - 0.765z) = 0$ and use bisection to solve it numerically, yielding $z_1 \approx 1.00295617$. Using proposition 9, we obtain $C \approx 1.042504$, so the probability that a read contains no seed is approximately $1.042504/1.00295617^{101} \approx 0.773749$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.77373 - 0.77376$.

★

Once again, the analytic combinatorics estimates are close to the exact values. Figure 11 illustrates the precision of the estimates for different values of the insertion rate r and of the read size k .

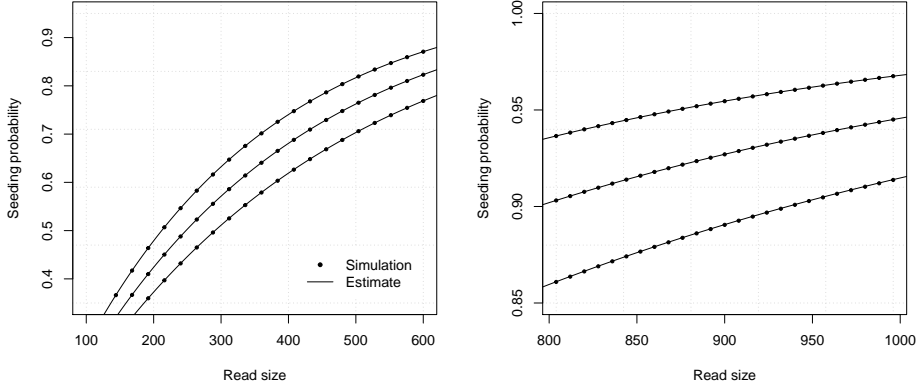


Figure 11: **Example estimates for substitutions, deletions and insertions.** The analytic combinatorics estimates of proposition 9 are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size contains a seed, either estimated by 10,000,000 random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$, $p = 0.05$, $\delta = 0.15$, $\tilde{r} = 0.45$ and $r = 0.04$, $r = 0.05$ or $r = 0.06$ (from top to bottom).

3.6 Empirical error models

In the theory developed so far, we introduced different kinds of errors because they have different probabilities and different sizes, but the nature of the error is irrelevant. To know whether the read contains a perfect seed, only their distribution matters.

An important consequence is that we can develop custom error models based on empirical estimates of the error distribution. This option is not only valid for modeling sequencing errors, but also for modeling mutations occurring through biological processes.

We introduce error-only intervals, which will encapsulate the available information about the size of error patches. Treating deletions separately will allow us to simplify the exposition, so we will consider that error-only intervals are always be non-empty.

Definition 7. An *error-only interval* is a non-empty sequence of errors that cannot be extended left or right.

We assume that every nucleotide has a constant probability q of being correct. With probability $p = 1 - q$, the nucleotide is incorrect and thus starts an error-only interval. If p_k is the empirical probability that the error-only interval has size k for $1 \leq k \leq n$, we can write the weighted generating function of error intervals as $pE(z)$, where $E(z) = p_1z + p_2z^2 + \dots + p_nz^n$.

The weighted generating function of deletions is denoted δ for consistency

with the previous sections. As before, we will ignore deletions adjacent to error-only intervals. Even if they occur, they have no consequence as they never interrupt a potential seed.

Reads under the empirical error model can be thought of as walks on the transfer graph shown in figure 12. An error-free interval can be followed by an error-only interval, or by another error-free interval if a deletion is present in between. An error-only interval can only be followed by an error-free interval.

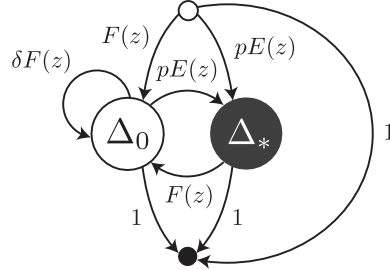


Figure 12: **Transfer graph of reads under the empirical error model.** Reads are viewed as sequences of error-free intervals (symbol Δ_0) or error-only intervals (symbol Δ_*). An error-free interval can follow another one if a deletion is present in between. $F(z)$ and $pE(z)$ are the weighted generating functions of error-free intervals and error-only intervals, respectively. $\delta F(z)$ is the weighted generating function of a deletion followed by an error-free interval.

The head and tail vectors are $(F(z), pE(z))^\top$ and $(1, 1)$, respectively; and $\psi(z) = 1$. The transfer matrix associated with the body of the transfer graph shown in figure 12 is

$$M(z) = \begin{bmatrix} \Delta_0 & \Delta_* \\ \Delta_0 & \Delta_* \end{bmatrix} \begin{bmatrix} \delta F(z) & pE(z) \\ F(z) & 0 \end{bmatrix}.$$

Computing $\psi(z) + H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$ as per proposition 4 yields

$$1 + [F(z) \ pE(z)] \cdot \frac{1}{1 - F(z)(\delta + pE(z))} \cdot \begin{bmatrix} 1 & pE(z) \\ F(z) & 1 - \delta F(z) \end{bmatrix} \cdot \begin{bmatrix} 1 \\ 1 \end{bmatrix},$$

which eventually simplifies to

$$R(z) = \frac{(1 + (1 - \delta)F(z))(1 + pE(z))}{1 - F(z)(\delta + pE(z))}. \quad (22)$$

As previously, the weighted generating function of error-free intervals is $F(z) = qz + (1 - \delta)(qz)^2 + (1 - \delta)^2(qz)^3 + \dots = qz/(1 - (1 - \delta)qz)$. Substituting this value in the equation above, we obtain

$$R(z) = \frac{1 + pE(z)}{1 - qz(1 + pE(z))}. \quad (23)$$

Expression (23) is in general not equal to $1/(1 - z)$ because $E(z)$ is not equal to $z/(1 - pz)$. Since the total weight of reads of size k is not equal to 1, we cannot simplify ratio (9) as in the previous cases. We will need to include $[z^k]R(z)$ in our calculations, which we will approximate with proposition 5. This implies that we have to find the root with smallest *modulus* of the polynomial $1 - qz(1 + \delta + E(z))$.

Remark 14. Why $[z^k]R(z) = 1$ for some models but not for others? The combinatorial “atoms” can be inhomogeneous, leading to a weight deficit for certain sequence sizes. For instance, sequences of the object aa with weighted generating function z^2 cannot have odd size. Empirical error-only intervals are typically bounded whereas error-free intervals are not. This imbalance introduces a weight deficit for some sequence sizes.

As in the previous sections, to find the weighted generating function of reads without an exact γ -seeds, we need to replace $F(z)$ in expression (22) by its truncation $F_\gamma(z) = qz + (1 - \delta)(qz)^2 + \dots + (1 - \delta)^{\gamma-2}(qz)^{\gamma-1}$. The terms only partially cancel each other and we obtain

$$S_\gamma(z) = \frac{(1 + pE(z))(1 - (1 - \delta)^\gamma(qz)^\gamma)}{1 - (1 - \delta)qz - qz(1 - (1 - \delta)^{\gamma-1}(qz)^{\gamma-1})(\delta + pE(z))}. \quad (24)$$

Remark 15. When $R(z)$ and $S_\gamma(z)$ are defined by (23) and (24), ratio (9) is always less than or equal to 1. To see this, note that expression (22) can be written as

$$\frac{1 + pE(z)}{\delta + pE(z)} (1 + (1 - \delta)F(z))(1 + F(z) + F(z)^2 + \dots). \quad (25)$$

From this expression, it is clear that replacing $F(z)$ by its truncation $F_\gamma(z)$ decreases the coefficient of z^γ (recall that all the weights are positive). This is equivalent to $[z^\gamma]R(z) \geq [z^\gamma]S_\gamma(z)$, confirming that the probability of occurrence of reads without seeds is always less than or equal to 1.

To estimate the total weight of seedless reads, we need to find the root with smallest modulus of $1 - qz(1 + E(z)) + (1 - \delta)^{\gamma-1}(qz)^{\gamma-1}(\delta + E(z))$. The solution depends on the particular expression of $E(z)$. Even though the process can be automated using proposition 5, every case is different and we cannot give an explicit formula here.

Example 17. Assume that empirical measurements suggest that $q = 0.9$, $\delta = 0.15$ and that error-only intervals of size 1, 2 or 3 have probabilities 0.5, 0.33 and 0.17, respectively (in this example, error-only intervals of size greater than 3 are never observed). This implies $E(z) = 0.5z + 0.33z^2 + 0.17z^3$.

If we choose seeds of size $\gamma = 17$, the weighted generating function of reads in expression (23) becomes

$$R(z) = \frac{1 + 0.1(0.5z + 0.33z^2 + 0.17z^3)}{1 - 0.9z(1 + 0.1(0.5z + 0.33z^2 + 0.17z^3))}.$$

The smallest root of the denominator is approximately equal to 1.008754. The multiplicative constant of proposition 5 is approximately equal to 0.962590, so the total weight of reads of size k is approximately equal to $0.962590/1.008754^{k+1}$.

Similary, the weighted generating function of reads without seeds from expression (24) becomes

$$S_{17}(z) = \frac{(1 + 0.1(0.5z + 0.33z^2 + 0.17z^3))(1 - (0.765z)^{17})}{1 - 0.765z - 0.9z(1 - (0.765z)^{16})(0.15 + 0.1(0.5z + 0.33z^2 + 0.17z^3))}.$$

The smallest root of the denominator is approximately 0.0122100. The ultiplicative constant of proposition 5 is approximately equal to 0.949377. So the total weight of seedless reads of size k is approximately equal to $0.949377/0.012098^{k+1}$.

Combining these two results, the probability that a read of size k has no seed is the ratio of the total weights computed above. This is approximately equal to $0.9862735/1.003315^{k+1}$.

*

3.7 Worst case for approximations

So far, all the examples showed that the analytic combinatorics approximations are accurate. Indeed, the main motivation for our approach is to find estimates that converge exponentially fast to the target value. Does this mean that we can always use the approximations in place of the true values?

To find out, we need to describe the behavior of the estimates in the worst conditions. The approximations become more accurate as the size of the sequence increases, *i.e.* as the reads become longer. This is somewhat awkward, as the read size is usually fixed by the technology or by the problem at hand. Still, the approximations described above tend to be less accurate for short reads.

The second aspect is convergence speed. In propositions 5 it was shown that the rate of convergence is dominated by the ratio between the two smallest singularities of the weighted generating function. This means that convergence is fastest when the dominant singularity is significantly closer to 0 than the other singularities. Conversely, convergence is slowest when at least one other singularity is almost as close to 0.

The worst case for the approximation is thus when the reads are small and when the parameters are such that singularities have relatively close *moduli*. In the error model of uniform substitutions, this corresponds to small values of the error rate p .

To see this, recall that the singularities $Re^{i\theta}$ of $S_\gamma(z)$ expressed in (12) are bound by the equation

$$p^2 R^2 = (pq^\gamma R^{\gamma+1})^2 + 2pR \cos(\theta) - 1. \quad (13)$$

Considering this an equation of $R > 0$, the solution is minimal when $2pR \cos(\theta)$ is maximal, and *vice versa*. Thus, the two equations for the smallest and largest solutions \underline{R} and \bar{R} are respectively

$$1 - p\underline{R} = pq^\gamma \underline{R}^{\gamma+1}, \text{ and}$$

$$1 + p\bar{R} = pq^\gamma \bar{R}^{\gamma+1}.$$

Observe that $\lim_{p \rightarrow 0} \underline{R} = \infty$, otherwise $z(1 + qz + \dots + (qz)^{\gamma-1})$ would be bounded and the equality $Q(z) = 1 - pz(1 + qz + \dots + (qz)^{\gamma-1}) = 0$ could not hold. This implies $\lim_{p \rightarrow 0} p\underline{R} = 0$ and $\lim_{p \rightarrow 0} p\bar{R} = 0$. Otherwise, if we had for instance $\lim_{p \rightarrow 0} p\underline{R} = \ell > 0$, taking the limit of the first equation above would yield $1 - \ell = \ell \underline{R}^\gamma$, inconsistent with the fact that \underline{R} is unbounded.

This last fact entails $\underline{R} \sim \bar{R} \sim \varphi(p) = 1/\sqrt[\gamma+1]{pq^\gamma}$. Indeed, we have $\underline{R}/\varphi(p) = (1 - p\underline{R})^{1/\gamma+1}$ and $\bar{R}/\varphi(p) = (1 + p\bar{R})^{1/\gamma+1}$, which both tend to 1 as p vanishes. This means that the *moduli* of all the singularities get closer to each other as p decreases. As a consequence, the speed of convergence of the approximation of proposition 5 diminishes (but it remains exponential).

In practical terms, the situation above describes the specifications of the Illumina technology, where errors are almost always substitutions, occurring at a frequency around 1% on current instruments. Since the reads are often around 50 nucleotides, the analytic combinatorics estimates of the seeding probabilities are typically less accurate than suggested in the previous sections.

Figure 13 shows the accuracy of the estimates in one of the worst cases. The curve is clearly distinct from the simulation at the chosen scale, but the absolute difference is never higher than approximately 0.01 (and lower for read sizes above 40). Whether this error is acceptable depends on the problem. Very often, p is known to within approximately 0.01, which is a more serious limitation on the precision. In most practical applications, the approximation error of proposition 5 can be tolerated even in the worst case, but it is important to bear in mind that it may not be totally negligible for reads of size 50 or lower.

If precision is crucial, remember that the exact solution can be found by using all the singularities of the weighted generating function S_γ (see remark 7). If computational speed is an issue, the singularities and associated proportionality constants can be precomputed for a range of values of p and γ .

3.8 Oversimplified error models

It is tempting to replace the somewhat complex error models of sections 3.4 and 3.5 by the uniform substitutions model of section 3.3 with an equivalent error rate. An intuitive approach would be to set the only parameter $q = 1 - p$ of this model to the value of $q(1 - \delta)$ of the more complex ones, because those represent the probability of decoding a nucleotide without error.

However, this approach is inaccurate because insertions and deletions can have a strong influence. To show this, let us revisit example 16 with an approximate substitution model instead of the full error model.

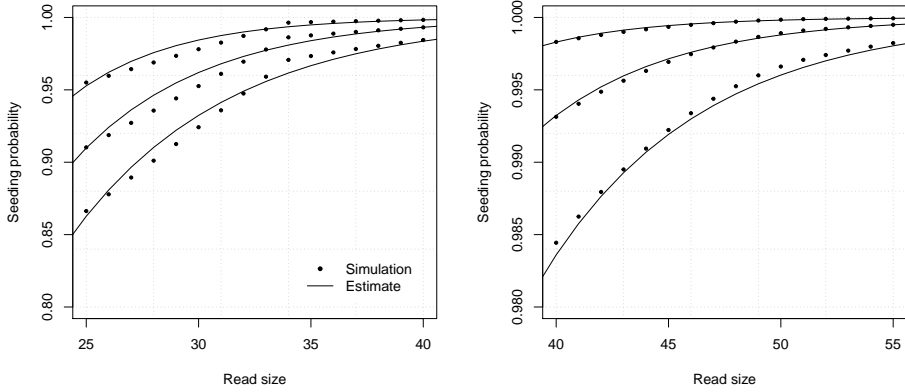


Figure 13: **Example worst case for seeding with substitutions only.** The analytic combinatorics estimates of proposition 7 are benchmarked against 10,000,000 random simulations. Shown on both panels are the probabilities that a read of given size contains a seed of size $d = 17$, either estimated by random simulations (dots), or by the estimate derived in section 3.3. The curves are drawn for $p = 0.005$, $p = 0.010$ or $p = 0.015$ (from top to bottom). The largest difference between the estimates and the simulations is around 0.01.

Example 18. In example 16, we computed the approximate probability that a read of size $k = 100$ contains no exact 17-seed, where $p = 0.05$, $\delta = 0.15$, $r = 0.05$ and $\tilde{r} = 0.45$. In such conditions, the probability of decoding a nucleotide without error (given that it is not in an insertion burst) is $(1 - p - r)(1 - \delta) = 0.765$. Now using a substitution model where $q = 0.765$ and thus $p = 0.235$, the analytic combinatorics estimate comes out as $1.03726 / 1.002591^{101} \approx 0.79866$. This number is outside the 99% confidence $0.77373 - 0.77376$ and the uniform substitution model underestimates the probability that the read contains a seed by more than 2 percentage points in this case.

*

The approximation error does not vanish. Actually, in the conditions of example 18 it increases with the read size k . Figure 14 shows the same data as figure 11, now fitted by an approximate substitution error. The error can be as high as 5 percentage points.

The conclusion is that the quality of the approximation collapses when using approximate error models. One may argue that an error of 5 percentage points may be tolerable, but the actual amount of error when using an approximate error model is unknown in general. Using simplified error models is unsafe because convergence is slow (the probability that the read contains a seed always tends to 1 as the size increases) and the error cannot be controlled.

Nevertheless, complexity comes at a cost. It is important to remember that the parameters of the error model may be inaccurate, in which case the

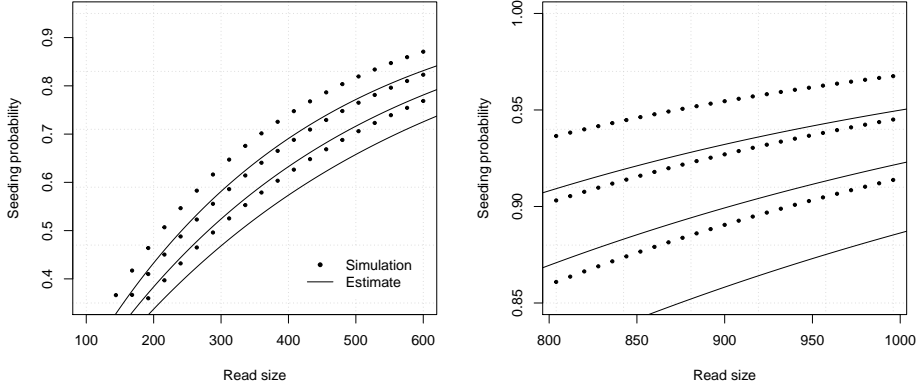


Figure 14: **Example estimates with oversimplified error model.** Shown on both panels are the probabilities that a read of given size contains a seed. The dots are obtained by 10,000,000 random simulations of the full error model with $\gamma = 17$, $p = 0.05$, $\delta = 0.15$, $\tilde{r} = 0.45$ and $r = 0.04$, $r = 0.05$ or $r = 0.06$ (from top to bottom); they are the same as in figure 14. The lines represent the analytic combinatorics estimates of the oversimplified uniform substitution model computed from proposition 7 with $p = 0.2265$ or $p = 0.2350$ or $p = 0.2435$ (top to bottom).

error is also not controlled. If these parameters are hard to estimate, it may be preferable to use a simplified model. But in general, the amount of data available in a sequencing run is sufficient to estimate the parameters of the error model. In practice, the safest approach to compute seeding probabilities is to use the full error model described in section 3.5 and estimate the four parameters from the alignment data generated during the mapping. If δ or r are small, they may be set to 0 for simplicity, yielding the error models of section 3.4 or section 3.3 (see remark 13).

4 Advanced seeding methods

Until now we have seen models of increasing complexity, but the transfer graphs had a fixed layout, and the weighted generating functions were ratios of polynomials of degree γ , the minimum size of the seed (with the notable exception of the empirical error model, for which the degree is not specified *a priori*). In relation to different problems, we will now explore other categories of models where the layout of the transfer graph depends on γ , and where the degree of the polynomials to solve will increase much faster than γ .

Because of this increased complexity, we will only consider the simplest uniform substitution error model. Otherwise, the strategy remains unchanged: we will formulate combinatorial problems, view their solutions as walks on transfer

graphs, encode these graphs as transfer matrices, obtain the weighted generating functions of the solutions from proposition 4 and approximate the answers using proposition 5.

4.1 Inexact seeds

We now consider a more challenging problem. The ongoing development of algorithms and data structures makes it possible to search inexact seeds, *i.e.* sequences that are very similar but not identical to the target. This comes at a greater computational cost than finding exactly similar sequences. If this cost is mitigated, it may be worth searching inexact seeds because the chances are higher to identify the target.

The theory below is implemented within the framework of the uniform substitution error model. Adaptations to more complex error models do not present new theoretical challenges, but the many cases to consider make the models cumbersome, at the cost of concision and clarity. Let us begin with a definition that will simplify the the following discussion.

Definition 8. A *single substitution interval* is a non-empty sequence of nucleotides that cannot be extended left or right, and that contains at most one substitution and no other error. An *inexact γ -seed* is a single substitution interval of size γ or greater.

We emphasize once more that our concern is the case of *one* substitution; the cases with more errors, or with errors of other types are not considered. An exact γ -seed is an inexact γ -seed, but the converse is not true. Figure 15 illustrates the relationship between substitutions and single substitution intervals. Note that single substitution intervals can overlap. In the models below, this is something that we will have to account for explicitly.

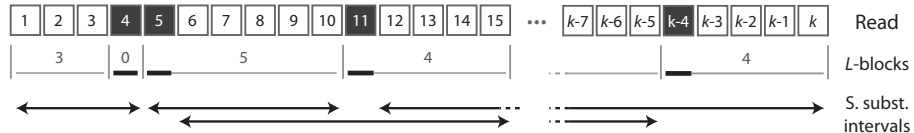


Figure 15: **Inexact seeding.** Substitutions (black squares) occur uniformly at random within the read. They delimit L -blocks and single substitution intervals. L -blocks consist of a substitution followed by an error-free interval. There is always a substitution on the left of an L -block, the error-free interval at the head of the read is not an L -block. Single substitution intervals (arrows) are the longest stretches of the read that contain at most one sbustitution.

We already computed the weighted generating function of reads under the uniform substitutions model in section 3.3. Namely, in equation (11) we found $R(z) = 1/(1 - z) = 1 + z + z^2 + \dots$ from which we concluded that the total weight of reads of size $k \geq 0$ is 1. We now need to find the weighted generating

function of reads that do not contain an inexact γ -seed. For this, we introduce a new type of combinatorial object.

Definition 9. An *L-block* is a substitution followed by an error-free interval.

The point of this definition is that a read is an error-free interval, followed by a sequence of *L-blocks*. In a read without inexact γ -seeds, not all the combinations of *L-blocks* are possible. For instance, an *L-block* with $\gamma - 2$ correct nucleotides can only be followed by a substitution, *i.e.* an *L-block* with 0 correct nucleotide (otherwise the concatenation of these two *L-blocks* forms an inexact γ seed). A substitution with $\gamma - 3$ correct nucleotides can be followed by an *L-block* with 0 or 1 correct nucleotide, *etc.*

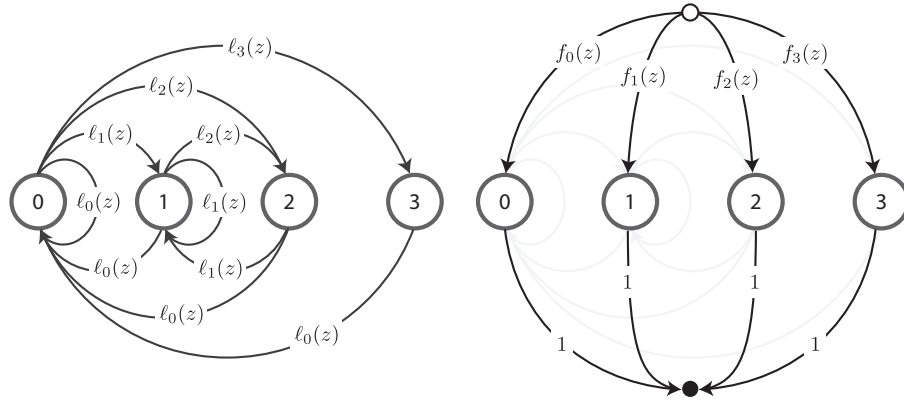


Figure 16: **Transfer graph of reads without inexact 5-seeds.** Reads are viewed as sequences of *L-blocks*. To not overload the figure, the body of the transfer graph is shown on the left, and the head and tail edges are shown on the right. The labels on the vertices represent the number of correct nucleotides in the *L-blocks*. The terms $\ell_i(z)$ are the weighted generating functions of *L-blocks* with i correct nucleotides for $i = 0, 1, 2, 3$. The terms $f_i(z)$ are the weighted generating functions of error-free intervals of size $i = 0, 1, 2, 3$.

To give a concrete example, reads without inexact 5-seeds can be seen as walks on the graph shown in figure 16. The numbers on the vertices indicates the amount of correct nucleotides in an *L-block*. There cannot be an *L-block* with 4 or more correct nucleotides, otherwise the read would contain an inexact 5-seed. The edges capture the relationships between consecutive *L-blocks*. The weighted generating function of *L-blocks* with i correct nucleotides is designated by $\ell_i(z)$, for $i = 0, 1, 2, 3$.

The head and tail vectors are shown on the the right part of figure 16. Reads can start with up to 3 correct nucleotides. The weighted generating function of error-free intervals of size i is designated $f_i(z)$, for $i = 0, 1, 2, 3$. Finally, any *L-block* can mark the end of the read, so all of them are connected to the tail vertex by 1, the weighted generating function of the the empty object ε . We do

not need to connect the head and the tail node with an such an edge, because as we will see below, the empty sequence is already described by the transfer graph.

The weighted generating function of an L -block with i correct nucleotides is $\ell_i(z) = pz \times qz \times \dots \times qz = pz(qz)^i$, i.e. the weighted generating function of a substitution followed by i correct nucleotides. In a similar way, the weighted generating function of a stretch of i correct nucleotides is $f_i(z) = qz \times \dots \times qz = (qz)^i$.

Because $f_0(z) = 1$, we now see that on the graph shown in figure 16, a walk starting from the head vertex and going to the tail vertex through the vertex labelled 0 has weighted generating function 1. This corresponds to the empty sequence, so there is no need to add it again to the graph, so $\psi(z) = 0$.

In the general case $\gamma > 0$, the matrix of the body of the transfer graph shown in figure 16 is

$$M(z) = \begin{bmatrix} 0 & 1 & 2 & \dots & \gamma-3 & \gamma-2 \\ \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ \gamma-4 & pz & pz(qz) & pz(qz)^2 & \dots & 0 \\ \gamma-3 & pz & pz(qz) & 0 & \dots & 0 \\ \gamma-2 & pz & 0 & 0 & \dots & 0 \end{bmatrix}.$$

The head vector $H(z)$ is equal to $(1, qz, (qz)^2, \dots, (qz)^{\gamma-2})^\top$, the tail vector $T(z)$ is equal to $(1, 1, \dots, 1)^\top$, and $\psi(z) = 0$. By proposition 4, the weighted generating function of reads without an inexact seed can be computed as $S_\gamma(z) = H(z) \cdot (I - M(z))^{-1} \cdot T(z)$.

Example 19. Let us revisit example 14, where we approximated the probability that a read of size $k = 100$ has no seed for $\gamma = 17$ and for $p = 0.1$; but this time we allow the seed to contain one substitution. The matrix $M(z)$ has dimension 16 and the weighted generating function $S_{17}(z)$ can be written as $P(z)/Q(z)$, where P and Q are polynomials. Using numerical methods to find the root of Q with smallest modulus, we obtain $z_1 \approx 1.079244$. Likewise, we obtain $-P(z_1)/Q'(z_1) \approx 2.326700$, so the probability that the read does not contain an inexact seed is approximately $2.326700/1.079244^{101} \approx 0.0010511$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.001049 - 0.001054$. In the same conditions, the chances that the read contains an exact seed was 90.4%, compared to 99.9% when one substitution is allowed.

*

The approximations have the typical accuracy of analytic combinatorics estimates. Figure 17 shows the precision of the estimates for different values of the substitution rate p and the read size k . Observe that the probability that

the read contains an inexact seed is substantially higher than the probability that it contains an exact seed (compare with figure 7). This means that efficient algorithms to find inexact seeds can greatly reduce the number of false negatives in the mapping problem.

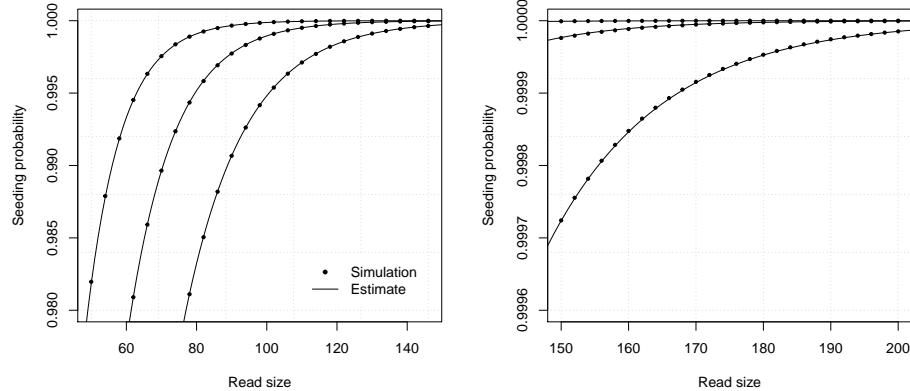


Figure 17: **Example estimates for inexact seeding with substitutions only.** The analytic combinatorics estimates described in this section are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size contains an inexact seed, either estimated by random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$, $p = 0.08$, $p = 0.10$ or $p = 0.12$ (from top to bottom). 10,000,000 simulations are run on the left panel and 100,000,000 on the right one.

Unfortunately, the expressions of the weighted generating functions given by proposition 4 are very cumbersome, and they may take hours to evaluate. For instance, the degrees of P and Q for $\gamma = 17$ are 152 and 153, respectively.

The best option here is to pre-compute the dominant singularities and associated multiplicative constants for a useful range of values of γ (up to 30 is sufficient for most applications) and of the parameter p (up to 0.25 is sufficient for most application). Once these values are stored, the estimates can be calculated rapidly.

4.2 Type I errors

Our concern so far was whether reads contain a seed, *i.e.* an error-free (or almost free) interval of sufficient size. If this is the case, the target sequence is among the candidates and we implicitly assume that it will be identified as the best hit during the alignment stage. But we did not discuss what happens if the read does not contain a seed.

There are two possible cases. The first is that the mapping procedure returns no hit. The conclusion is then that the sequence does not belong to the genome,

or that it cannot be mapped. In analogy with statistical testing, we will call this a “type II error”. The second case is that the mapping procedure returns a hit. This cannot be a true hit because the target was discarded during the seeding stage. We will call this a “type I error”. Type I errors are the most difficult to detect, because nothing distinguishes them from true hits, at least at the seeding stage.

It is important to emphasize that in this version of the mapping problem, the split between type I and type II errors amounts to the probability that the read contains no seed (because we assume that all the candidates are aligned and that the alignment always identifies the true hit when it is present among the candidates). This split depends on the genome, and more particularly on the configuration of its repeated sequences. A stretch of γ nucleotides matches a random genome with probability $O(4^{-\gamma})$. However, genomes are not a succession of random nucleotides. Large sequences are often duplicated, to the extent that the greater part of eukaryotic genomes can be considered repetitive. In addition, repeats are usually not exactly identical, but only similar. This means that the $O(4^{-\gamma})$ term can be a spectacular underestimate.

A complete treatment of type I errors is presently not possible. We will focus on a single case that will be the basis for further development. More specifically, we will assume that the target sequence has exactly one duplicate in the genome, and that they differ from each other only by uniform substitutions (we will see below how to relax these assumptions). The problem then amounts to finding the probability that the seeding process identifies the duplicate as a candidate, but not the target sequence. Figure 18 shows how this can occur.

Error		
	ATGCGAGTAGCGCTG A GGAGAGCGTATTAA	Read
	ATGCGAGTAGCGCTG T GGAGAGCGTATTAA	Target
	A CACGAGTAGCGCTG A GGAGAGCGTATTAAA	Duplicate

Figure 18: **Example type I error.** A read of 30 nucleotides contains a single error at the sixteenth position. The erroneous nucleotide is identical to the duplicate sequence at this position. This creates a match of 24 nucleotides for the duplicate sequence, while the longest match for the target is 15 nucleotides. Thus, using seeds of size greater than 15 would result in a type I error (even though the duplicate has three differences with the read and the target has only one).

Remark 16. *With only one reference sequence, “error-free” and “matching” nucleotides are the same. With more than one reference, we must take care of distinguishing them. In this case, matching nucleotides can be incorrect (they can match the wrong sequence).*

Instead of tackling this problem directly, we will set a more general framework where a read can match two similar sequences denoted (+) and (-). A read now consists of four types of nucleotides: those matching both sequences, those matching the (+) sequence only, those matching the (-) sequence only,

and those matching none. The first type will be referred to as a *match* and the last three as *mismatches* (single or double). The following definitions will be useful to sketch the transfer graph.

Definition 10. An **R -block** is a mismatch-free interval followed by a single or a double mismatch. A (+) **interval** (respectively (-) **interval**) is a sequence of nucleotides matching the (+) sequence (respectively (-) sequence) that cannot be extended left or right. A (+) **seed** (respectively (-) **seed**) is a (+) interval (respectively (-) interval) of size γ or greater.

This information is summarized in figure 19. We will use it to find a construction for reads without either (+) or (-) seeds. We will later show how this can be used to find the probabilities of type I errors.

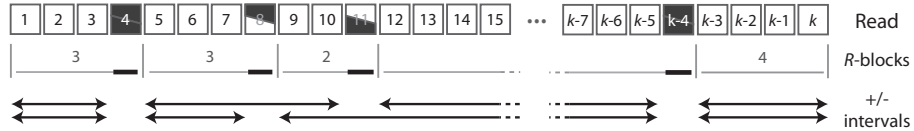
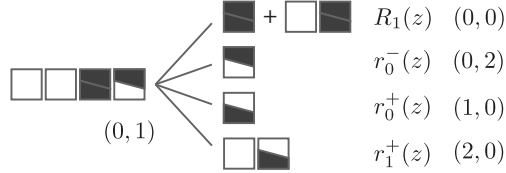


Figure 19: **Seeding with two reference sequences.** Matches are represented as white squares, mismatches against the (+) sequence as bottom black wedges, mismatches against the (-) sequence as top black wedges, and double mismatches as black squares. They delimit R -blocks consisting of mismatch-free intervals followed by a mismatch. The last nucleotide of an R -block is always a mismatch so the mismatch-free interval at the tail of the read is not an R -block. (+) or (-) intervals are stretches of the read that match either the (+) sequence (top arrow) or the (-) sequence (bottom arrow) and that cannot be extended left or right.

Remark 17. When a subsequence is both a (+) and a (-) seed (the head and the tail of the read shown in figure 19 for instance), then both sequences are candidates.

Reads can be viewed as sequences of R -blocks followed by a mismatch-free interval. Reads without (+) or (-) seeds have particularly complex constraints on the arrangements of those R -blocks. For instance, an R -block of size γ (*i.e.* $\gamma - 1$ matches followed by a mismatch) must be followed by a mismatch if the final nucleotide is a single mismatch, but it can be followed by up to $\gamma - 1$ matches if the last nucleotide is a double mismatch.

To give a concrete example, we will consider reads without (+) or (-) seeds of size 3. The read below features two matches, followed by a double mismatch, followed by a mismatch against the (-) sequence. The numbers between brackets (0, 1) indicate the respective amount of nucleotides matching the (-) and the (+) sequences at the end of the R -block. At that point, there are only five possibilities for the next R -block; every other combination would create a (+) seed or a (-) seed.



Appending a double mismatch or a match followed by a double mismatch (top case) brings the number of nucleotides matching $(-)$ and $(+)$ at the end of the R -block to $(0, 0)$. The weighted generating function of this union of R -blocks is denoted $R_1(z)$. Appending a mismatch against the $(-)$ sequence (second case from the top), *i.e.* an R -block with weighted generating function $r_0^-(z)$ brings the amount of matching nucleotides to $(0, 2)$. Conversely, appending a mismatch against the $(+)$ sequence (third case from the top), *i.e.* an R -block with weighted generating function $r_0^+(z)$ brings the amount of matching nucleotides to $(1, 0)$. Finally, appending a match followed by a mismatch against the $(+)$ sequence (bottom case), *i.e.* an R -block with weighted generating function $r_1^+(z)$ brings the amount of matching nucleotides to $(2, 0)$.

By considering all the cases, we obtain the transfer graph shown in figure 20. Even for such a small value of γ , the graph is so dense that the edges have been separated in two panels and encoded as symbols in order to not overload the figure.

As in the example above, the numbers in the vertices indicate the amount of nucleotides matching $(-)$ and $(+)$ at the end of the R -blocks. Since R -blocks are terminated by a mismatch, at least one of these numbers is 0. The body of the transfer graph is only partially represented on the left panel. The edges pointing to the vertex $(0, 0)$ are displayed on the right panel, together with the head and tail edges. The four kinds of edges on the left panel correspond to different R -blocks with weighted generating functions $r_0^+(z)$, $r_0^-(z)$, $r_1^+(z)$ and $r_1^-(z)$. The sign indicates which sequence is *mismatched* at the end of the R -block and the index is the number of matches in the R -block.

The edges of the body of the transfer graph on the right panel point to the vertex $(0, 0)$. All the vertices are connected to this vertex because appending an R -block terminated by a double mismatch always brings the read to this state. The weighted generating functions $R_0(z)$, $R_1(z)$ and $R_2(z)$ represent such R -blocks with *up to* 0, 1 or 2 matches, respectively.

In the initial state of the read, the amount of matching nucleotides is $(0, 0)$, which is indicated by the label 1 on the edge between the head vertex and the vertex $(0, 0)$. This is equivalent to prepending the read by the empty object ε . Reads are terminated by a sequence of matches. The weighted generating functions $F_0(z)$, $F_1(z)$ and $F_2(z)$ represent mismatch-free intervals of size *up to* 0, 1 or 2, respectively. Here $\psi(z) = 0$. The empty sequence is present in the graph (take a path from the head vertex to the tail vertex through $(0, 0)$ with no match appended at the tail), and no extra sequence needs to be added.

The same logic can be applied for higher values of γ . The transfer graph

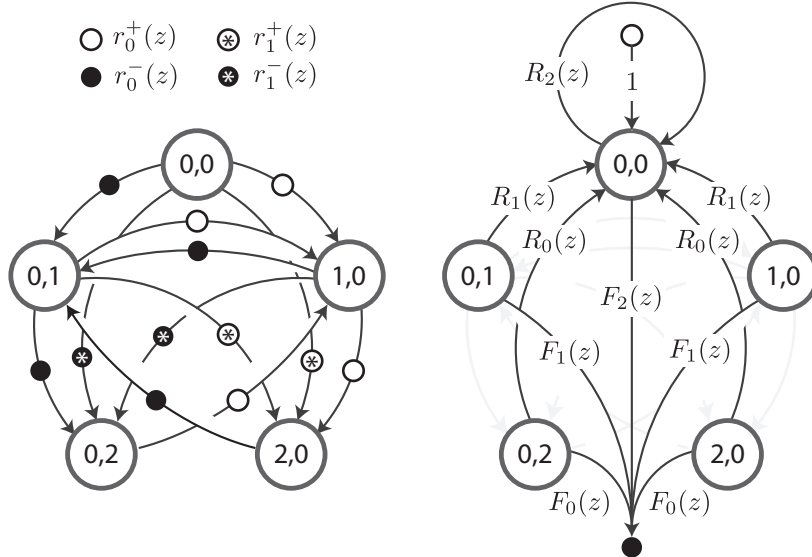


Figure 20: **Transfer graph of reads without seed of size 3 for either (+) or (-).** Reads are viewed as sequences of R -blocks, and they can match two sequences referred to as (+) and (-). To not overload the figure, the edges are split between the left and right panels. The labels on the vertices represent the respective number of nucleotides matching the (-) and (+) sequences at the end of an R -block. Weighted generating functions $r_i^+(z)$ represent R -blocks with i matches terminated by a mismatch against the (+) sequence (symmetrically for the (-) sequence). Weighted generating function $R_i(z)$ represent R -blocks with *up to* i matches terminated by a double mismatch. Weighted generating functions $F_i(z)$ represent mismatch-free intervals of *up to* i matches.

becomes too cumbersome to represent but the transfer matrix has enough regularity to be specified in full. The general transfer matrix associated with the body of the transfer graph has dimension $2\gamma - 1$ and is defined as

$$M(z) = \begin{bmatrix} & & & & & & \\ & 0,0 & 1,0 & \cdots & \gamma-1,0 & 0,1 & \cdots & 1,\gamma-1 \\ & R_{\gamma-1}(z) & r_0^+(z) & \cdots & r_{\gamma-2}^+(z) & r_0^-(z) & \cdots & r_{\gamma-2}^-(z) \\ & 1,0 & R_{\gamma-2}(z) & & & & & \\ & 2,0 & R_{\gamma-3}(z) & & & & & \\ & \vdots & \vdots & & A(z) & & B(z) & \\ & \gamma-1,0 & R_0(z) & & & & & \\ & 0,1 & R_{\gamma-1}(z) & & & & & \\ & 0,2 & R_{\gamma-2}(z) & & & & & \\ & \vdots & \vdots & & C(z) & & D(z) & \\ & 0,\gamma-1 & R_0(z) & & & & & \end{bmatrix},$$

where $A(z)$, $B(z)$, $C(z)$ and $D(z)$ are square matrices with dimension $\gamma - 1$ and defined as

$$A(z) = \begin{bmatrix} & 1,0 & 2,0 & \cdots & \gamma-2,0 & \gamma-1,0 \\ & 0 & r_0^+(z) & \cdots & r_{\gamma-3}^+(z) & r_{\gamma-2}^+(z) \\ & 2,0 & 0 & \cdots & r_{\gamma-4}^+(z) & r_{\gamma-3}^+(z) \\ & \vdots & \vdots & \ddots & \vdots & \vdots \\ & \gamma-2,0 & 0 & 0 & \cdots & 0 & r_0^+(z) \\ & \gamma-1,0 & 0 & 0 & \cdots & 0 & 0 \end{bmatrix},$$

$$B(z) = \begin{bmatrix} & 0,1 & 0,2 & \cdots & 0,\gamma-2 & 0,\gamma-1 \\ & r_0^-(z) & r_1^-(z) & \cdots & r_{\gamma-3}^-(z) & r_{\gamma-2}^-(z) \\ & 2,0 & r_0^-(z) & r_1^-(z) & \cdots & r_{\gamma-3}^-(z) & 0 \\ & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ & \gamma-2,0 & r_0^-(z) & r_1^-(z) & \cdots & 0 & 0 \\ & \gamma-1,0 & r_0^-(z) & 0 & \cdots & 0 & 0 \end{bmatrix},$$

$$C(z) = \begin{bmatrix} & 1,0 & 2,0 & \cdots & \gamma-2,0 & \gamma-1,0 \\ & r_0^+(z) & r_1^+(z) & \cdots & r_{\gamma-3}^+(z) & r_{\gamma-2}^+(z) \\ & 0,1 & r_0^+(z) & r_1^+(z) & \cdots & r_{\gamma-3}^+(z) & 0 \\ & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ & 0,\gamma-2 & r_0^+(z) & r_1^+(z) & \cdots & 0 & 0 \\ & 0,\gamma-1 & r_0^+(z) & 0 & \cdots & 0 & 0 \end{bmatrix},$$

$$D(z) = \begin{bmatrix} & 0,1 & 0,2 & \dots & 0,\gamma-2 & 0,\gamma-1 \\ 0,1 & 0 & r_0^-(z) & \dots & r_{\gamma-3}^-(z) & r_{\gamma-2}^-(z) \\ 0,2 & 0 & 0 & \dots & r_{\gamma-4}^-(z) & r_{\gamma-3}^-(z) \\ \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ 0,\gamma-2 & 0 & 0 & \dots & 0 & r_0^-(z) \\ 0,\gamma-1 & 0 & 0 & \dots & 0 & 0 \end{bmatrix}.$$

As per proposition 4, the weighted generating function of reads with no (+) nor (−) seeds is given by $H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$. In this case the head vector $H(z)$ is equal to $(1, 0, 0, \dots, 0)^\top$ and the tail vector $T(z)$ is equal to $(R_{\gamma-1}(z), R_{\gamma-2}(z), \dots, R_0(z), R_{\gamma-2}(z), \dots, R_0(z))^\top$.

Under the assumption that mismatches occur uniformly in the read, nucleotides of each type occur with probabilities a, b, c or d , with $a + b + c + d = 1$, where a is the probability that the nucleotide is a double match, b that it is a mismatch against (+), c that it is a mismatch against (−) and d that it is a mismatch against both. With these notations we obtain

$$\begin{aligned} r_i^+(z) &= (az)^i cz, \\ r_i^-(z) &= (az)^i bz, \\ R_i(z) &= (1 + az + \dots + (az)^i) dz, \\ F_i(z) &= 1 + az + \dots + (az)^i. \end{aligned} \tag{26}$$

We could now find the weighted generating function of the reads without (+) or (−) seeds, but we will return to the original problem of computing the probability of a type I error. In this context, the (+) sequence is the target and the (−) sequence is the duplicate.

We will assume that each nucleotide is decoded incorrectly with probability p and that for each nucleotide, the duplicate sequence differs from the target with probability κ . A nucleotide is a mismatch for both sequences if it is a read error (probability p) and if either the duplicate is identical to the target (probability $1 - \kappa$) or it is different from both the target and the decoded nucleotide (probability $2\kappa/3$). The other probabilities can be computed with similar arguments, yielding $a = (1 - p)(1 - \kappa)$, $b = p\kappa/3$, $c = (1 - p)\kappa$ and $d = p(1 - \kappa)/3$.

The model is completely specified by p and κ . With their actual values at hand, we can use proposition 5 to obtain an asymptotic estimate of the probability that a read has no match for the target *and* no match for the duplicate. Call this estimated probability P_2 . Using the results of section 3.3, we can also compute an asymptotic estimate of the probability that the read does not contain an exact γ -seed (irrespective of potential matches to the duplicate). Call this estimated probability P_1 . We can thus estimate the probability that the read contains a match of size γ or greater for the duplicate sequence *but* no exact γ -seed as $P_1 - P_2$.

Example 20. Let us approximate the probability of type I error (with one duplicate) for a read of size $k = 100$ with $\gamma = 17$ and for substitution rates $p = 0.10$ and $\kappa = 0.10$. In example 14 we found $P_1 \approx 1.396145/1.0268856^{101}$. To compute P_2 , we substitute $a = 0.81, b = 0.09, c = 0.00333, d = 0.09667$ in the expression of the 33×33 transfer matrix $M(z)$ and in the expression of $T(z)$. We then compute the expression $H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$ and obtain a function $P(z)/Q(z)$ where P and Q are polynomials of degree 288 and 289, respectively. The root of Q with smallest modulus is the dominant singularity z_1 of the weighted generating function. Using numerical approaches, we find $z_1 \approx 1.0272930245$. We compute the proportionality constant of proposition 5 as $P(z_1)/Q'(z_1) \approx 1.4032791$ from which we obtain $P_2 \approx 1.4032791/1.0272930245^{101}$. The type I error rate is then approximately equal to 0.0032906. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.003288 - 0.003292$.

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Figure 21 illustrates the precision of this estimate for different values of the substitution rate κ and of the read size k . Observe that the estimates are accurate even for such low probability of occurrence.

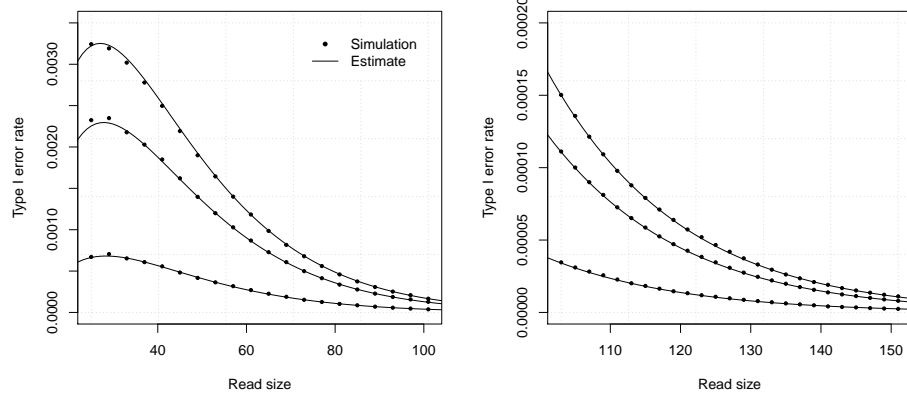


Figure 21: **Example estimates of type I errors (one duplicate).** The analytic combinatorics estimates are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size will give a type I error, either estimated by 10,000,000 (left) or 100,000,000 (right) random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$ and $p = 0.05$, $\kappa = 0.05, \kappa = 0.15$ or $\kappa = 0.25$ (from top to bottom).

Computing P_2 with proposition 5 requires finding a weighted generating function $P(z)/Q(z)$ where P and Q are polynomials with too many terms to fit in this document. It is thus problematic to compute the approximations efficiently. The best option is once again to pre-compute the dominant singularities and associated multiplicative constants for a useful range of γ (up to 30

is sufficient for most applications) and of the parameters p and κ (up to 0.25 is sufficient for most applications). Once these values are stored, the estimates can be calculated rapidly without having to compute the weighted generating function and finding its dominant singularity.

We have started this section with the assumption that the target has exactly one duplicate. Also, κ is usually unknown so how are the estimates derived above useful?

First, it is important to stress that the seeding strategy is efficient only when the target has few duplicates. Some sequences have more than 100,000 copies in the human genome. In such cases, the seeding process yields many candidates that will cost time at the alignment stage, for in the end relatively small chances of identifying the correct target. As a consequence, most mapping algorithms bail out as soon as they gather enough evidence that the target has many duplicates. There is thus little incentive to develop accurate estimates of type I error for highly repeated sequences, and one should focus on the case of intermediate amount of repeats, say around 20.

We can extend the theory developed above to more than one duplicate. The probability that a read contains no match for the duplicate given that it contains no match for the target is P_2/P_1 . If the target has N extra duplicates evolving independently with mutation rate κ , the probability that a read contains no match for any of the duplicates given that it contains no match for the target is $(P_2/P_1)^N$. The probability of a type I error is thus $P_1(1 - (P_2/P_1)^N)$. For $N = 1$, we recover $P_1 - P_2$ and for large N , the value approaches P_1 , the probability that the read has no seed.

This approach is somewhat naive because repeated sequences do not evolve independently of each other and N is usually unknown. However, it gives a handle on the problem as N and κ can be estimated during the seeding or better, stored for every position of the genome.

Also observe that the case of one duplicate gives a lower bound for the probability of type I error. When there is exactly one duplicate, type I errors are almost impossible to detect because the mapping process identifies a single hit that is similar to the read (as if a seed was present and the target had no duplicate). If the proportion of the genome that is repeated is x and P^* is the maximum of $P_1 - P_2$ relative to unknown κ , then xP^* is a reasonable lower bound on the probability of type I errors. If more information about the read, or about the seeding and alignment processes is available, then this lower bound can vary.

4.3 MEM seeds

In the exact seeding scheme considered so far, every subsequence of the genome that matches γ consecutive nucleotides of the read is considered a candidate. While assuring good chances of discovering the target, this approach often yields “too many” useless candidates, which costs computational time at the alignment stage. An alternative scheme called MEM seeding (Maximal Exact Match) gives

better empirical results. The principle is to use as seeds only the longest local matches between the read and the genome.

Definition 11. A γ -MEM (Maximal Exact Match) is a sequence of at least γ nucleotides from the read, that matches a subsequence of the genome and that cannot be extended left or right.

Remark 18. Note that error-free intervals and mismatch-free intervals are disjoint, whereas MEMs can be overlapping (and usually are).

Since there are fewer MEM seeds than exact γ -seeds, MEM seeding is more prone to errors than exact seeding. As we will see below, an important difference is that the combined frequencies of type I and type II errors is strictly greater than the probability that the read contains an error-free interval of size γ or greater.

To make the problem more concrete, we will consider as in section 4.2 that the target has exactly one duplicate sequence. In a similar way, we will start by defining two potential matches referred to as (+) and (-). The definitions of R -block, (+) interval and (-) interval given in section 4.2 still apply.

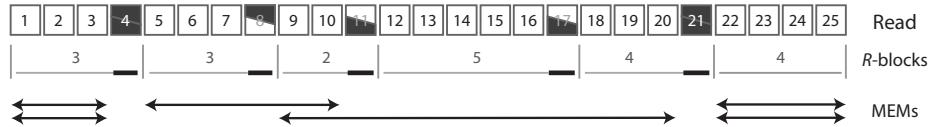


Figure 22: **MEMs (with two references)**. The symbols are the same as in figure 19. Matches are represented as white squares, mismatches for the (+) sequence as bottom black wedges, mismatches for the (-) sequence as top black wedges, and double mismatches as black squares. They delimit R -blocks consisting of mismatch-free intervals followed by a mismatch. The last nucleotide of an R -block is always a mismatch so the mismatch-free interval at the tail of the read is not an R -block. MEMs (arrows) are stretches of the read that match any of the two sequences and that cannot be extended left or right. MEMs matching the (+) sequence are represented at the top and MEMs matching the (-) sequence at the bottom. In this example, the central stretch of 5 nucleotides matching the (+) sequence is not used as seed because it is included in a stretch of 12 nucleotides matching the (-) sequence.

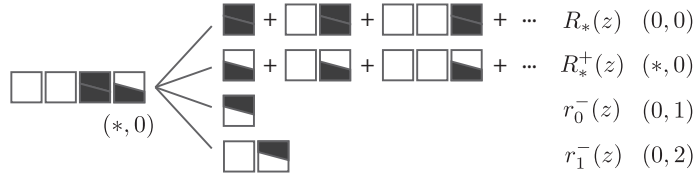
Figure 22 highlights the properties of MEMs. Notice how a match for (+) is masked by a longer match for (-). Nucleotides from 12 to 16 match the (+) sequence, creating a potential seed. However, it is contained in a match for the (-) sequence from nucleotides 9 to 20. Since the match for (+) is not a MEM, it is not used as a seed. Cases such as this one can cause type I errors to occur even when the reads contains matches of size γ or greater for the target.

Remark 19. When a MEM matches both sequences (the head and the tail of the read shown in figure 22 for instance), then both sequences are candidates.

As in section 4.2, reads are seen as sequences of R -blocks. We will first focus on the probability that a read does not contain a γ -MEM seed for the (+)

sequence, from which we will later deduce the probabilities of type I and type II errors. This restriction imposes relatively complex constraints on the R -blocks. For instance, an R -block of size γ (*i.e.* $\gamma - 1$ matches followed by a mismatch) must be followed by a mismatch if the final nucleotide is a mismatch for the $(-)$ sequence, but it may be followed by up to $\gamma - 1$ matches if this nucleotide is a double mismatch, or by any number of matches if it is a mismatch for the $(+)$ sequence.

To give a concrete example, we will consider γ -MEMs of size 3. The read below features two matches, followed by a double mismatch, followed by a mismatch for the $(+)$ sequence. The symbols between brackets $(*, 0)$ indicate the respective amount of nucleotides matching the $(-)$ and the $(+)$ sequences at the end of the R -block. All the R -blocks terminated by a mismatch against the $(+)$ sequence are equivalent because the subsequent R -block cannot be part of a MEM for the $(+)$ sequence. So the only information we need regarding the $(-)$ sequence is whether this amount is greater than 0, which we represent by the $*$ symbol. At that point, there are infinitely many possibilities for the next R -blocks, but only four distinct classes do not create a MEM seed for the $(+)$ sequence.



Appending any number of matches followed by a double mismatch (top case) brings the number of nucleotides matching $(-)$ and $(+)$ at the end of the R -block to $(0, 0)$. The weighted generating function of this union of R -blocks is denoted $R_*(z)$. Appending any number of matches followed by a mismatch against the $(+)$ sequence (second case from the top) maintains the amount of matching nucleotides as $(* , 0)$. The weighted generating function of this union of R -blocks is denoted $R_*^+(z)$. Appending a mismatch for the $(-)$ sequence (third case from the top), *i.e.* an R -block with weighted generating function $r_0^-(z)$ brings the amount of matching nucleotides to $(0, 1)$. Finally, appending a match followed by a mismatch for the $(-)$ sequence (bottom case), *i.e.* an R -block with weighted generating function $r_1^-(z)$ brings the amount of matching nucleotides to $(0, 2)$.

By considering all the cases, we obtain the transfer graph shown in figure 23. The numbers in the vertices indicate the amount of nucleotides matching $(-)$ and $(+)$ at the end of the R -blocks. Since R -blocks are terminated by a mismatch, at least one of these numbers is 0. As explained above, the $*$ symbol stands for one or more nucleotides, as all these cases are equivalent. The body of the transfer graph is represented on the left panel, the head and tail edges on the right panel.

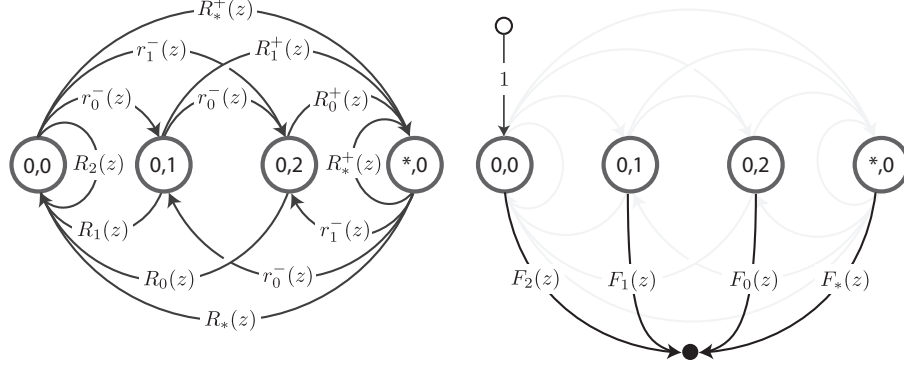


Figure 23: **Transfer graph of reads without 3-MEM seed for (+).** Text.

Simple R -blocks have weighted generating functions $r_0^+(z)$, $r_0^-(z)$, $r_1^+(z)$ or $r_1^-(z)$. The sign indicates which sequence is *mismatched* at the end of the R -block and the index indicates the number of matches in the R -block. Unions of R -blocks have weighted generating functions $R_*(z)$, $R_*^+(z)$ or $R_i^+(z)$. The first represents any number of matches followed by a double mismatch, the second represents any number of matches followed by a mismatch against the (+) sequence, the last represent *up to* i matches followed by a mismatch against the (+) sequence.

In the initial state of the read, the amount of matching nucleotides is $(0,0)$, which is indicated by the label 1 on the edge between the head vertex and the vertex $(0,0)$. This is equivalent to prepending the read by the empty object ε . Reads are terminated by a sequence of matches. The weighted generating functions $F_*(z)$, $F_0(z)$, $F_1(z)$ and $F_2(z)$ represent mismatch-free intervals of any size or size *up to* 0, 1 or 2, respectively. Here $\psi(z) = 0$. The empty sequence is present in the graph (take a path from the head vertex to the tail vertex through $(0,0)$ with no match appended at the tail), and no extra sequence needs to be added.

The same logic can be applied for higher values of γ . The general transfer matrix associated with the body of the transfer graph has dimension $\gamma + 1$ and is defined as

$$M(z) = \begin{bmatrix} & 0,0 & 0,1 & 0,2 & \dots & 0,\gamma-1 & *,0 \\ 0,0 & R_{\gamma-1}(z) & r_0^-(z) & r_1^-(z) & \dots & r_{\gamma-2}^-(z) & R_*^+(z) \\ 0,1 & R_{\gamma-2}(z) & 0 & r_0^-(z) & \dots & r_{\gamma-3}^-(z) & R_{\gamma-2}^+(z) \\ 0,2 & R_{\gamma-3}(z) & 0 & 0 & \dots & r_{\gamma-4}^-(z) & R_{\gamma-3}^+(z) \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots \\ 0,\gamma-1 & R_0(z) & 0 & 0 & \dots & 0 & R_0^+(z) \\ *,0 & R_*(z) & r_0^-(z) & r_1^-(z) & \dots & r_{\gamma-2}^-(z) & R_*^+(z) \end{bmatrix}.$$

As per proposition 4, the weighted generating function of reads with no γ -MEM for the (+) sequence is given by $H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$. In this case the head vector $H(z)$ is equal to $(1, 0, \dots, 0)^\top$ and the tail vector $T(z)$ is equal to $(F_{\gamma-1}(z), \dots, F_1(z), F_0(z), F_*(z))^\top$.

As in section 4.2, denote a the probability that the nucleotide is a double match, b that it is a mismatch against (+), c that it is a mismatch against (-) and d that it is a mismatch for both. With these definitions, expressions (26) still apply, and we also have

$$\begin{aligned} R_*(z) &= (1 + az + (az)^2 + \dots)dz = \frac{dz}{1 - az}, \\ R_*^+(z) &= (1 + az + (az)^2 + \dots)c z = \frac{cz}{1 - az}, \\ R_i^+(z) &= (1 + az + \dots + (az)^i)c z, \\ F_*(z) &= 1 + az + (az)^2 + \dots = \frac{1}{1 - az}. \end{aligned}$$

Knowing the values of a , b , c and d , we can compute the weighted generating function $H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$ and use proposition 5 to obtain an asymptotic estimate of the probability that a read does not contain a γ -MEM for the (+) sequence.

We now return to the problem of finding the probability that a read contains a γ -MEM seed when the target sequence has exactly one duplicate. If we consider that the (+) sequence is the target and that the (-) sequence is the duplicate, the probability described above is the one we are searching. The model is completely specified by the substitution rates p and κ , and as in section 4.2 $a = (1 - p)(1 - \kappa)$, $b = p\kappa/3$, $c = (1 - p)\kappa$ and $d = p(1 - \kappa)/3$. The example below shows how the computations are carried out in practice.

Example 21. Let us approximate the probability that a read does not contain a MEM seed (where the target has one duplicate) for a read of size $k = 100$ with $\gamma = 17$ and for $p = 0.10$ and $\kappa = 0.10$. First, we substitute in the 18×18 transfer matrix $M(z)$ and in $T(z)$ the values $a = 0.81$, $b = 0.09$, $c = 0.00333$, $d = 0.09667$. We then compute the expression $H(z)^\top \cdot (I - M(z))^{-1} \cdot T(z)$ and obtain a function $P(z)/Q(z)$ where P and Q are polynomials of degree 35 and 36, respectively. The root of Q with smallest modulus is the dominant singularity z_1 of the weighted generating function. Using numerical approaches, we find $z_1 \approx 1.0266331946$. We compute the proportionality constant of proposition 5 as $P(z_1)/Q'(z_1) \approx 1.3893346$ from which we obtain the final estimate $1.3893346/1.0266331946^{101} \approx 0.09769252$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.097682 - 0.097698$.

*

Figure 24 illustrates the precision of this estimate for different values of the error rate p and of the read size k . The values of the parameters are chosen to match those of figure 7 where we used exact seeds. In the conditions above, the seeding probabilities are almost indistinguishable. This does not mean that the cost of using MEM seeds is always negligible. When using MEMs, the seeding

probability goes down as the number of duplicates goes up (here we assume that there is only one duplicate). In addition we are not separating type I and type II errors, which typically have different costs.

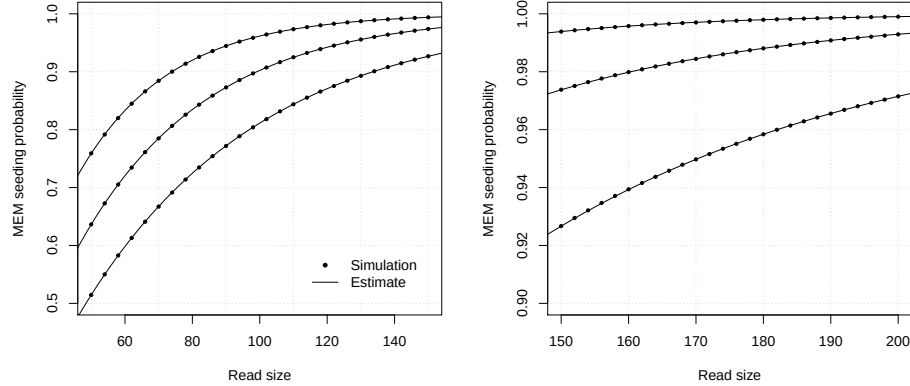


Figure 24: **Example estimates of MEM seeding probability (one duplicate).** The analytic combinatorics estimates are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size will contain a MEM seed, either estimated by 10,000,000 random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$ and $\kappa = 0.10$, $p = 0.08$, $p = 0.10$ or $p = 0.12$ (from top to bottom). Notice how close the values are to those shown in figure 7 (probabilities that the read contains an exact seed with the same parameter values).

This raises the question of how to compute the type I error rate when using MEM seeds. The work above provides all the elements we need to approximate it. Call the P_3 the approximate probability that the read does not contain a γ -MEM seed. Using the results from section 4.2, we can compute an asymptotic estimate of the probability that a read does not contain an exact γ -seed *and* does not contain any stretch of size γ or greater matching the duplicate sequence. We previously called this estimated probability P_2 . We can estimate the probability of type I errors as $P_3 - P_2$.

Let us clarify the last statement. Call A the event that the read has no γ -MEM seed and B the event that it does not contain an exact γ -seed *and* does not contain any stretch of size γ or greater matching the duplicate sequence. First observe that $B \subset A$ so $P(A \cap B^c) = P(A) - P(B)$. Then, B^c is the event that the read contains an exact γ -seed, a stretch of size γ or greater matching the duplicate sequence, or both. Another way to describe B^c is that “some hit is found in the read”. So $A \cap B^c$ is the event that “some hit is found, but it is not the target”, which is the definition of a type I error. Since $P(A) \approx P_3$ and $P(B) \approx P_2$, the probability of type I errors is indeed approximately equal to $P_3 - P_2$.

Remark 20. The rationale above shows that the probability of type II error with MEM seeds is approximately equal to the probability P_2 computed at section 4.2.

The example below shows the probabilities of type I errors are computed in practice.

Example 22. Let us approximate the probability of type I error with MEM seeds (with one duplicate) for a read of size $k = 100$ with $\gamma = 17$ and for substitution rates $p = 0.10$ and $\kappa = 0.10$. In example 20 we found $P_2 \approx 1.4032791/1.0272930245^{101}$ and in example 21 we found $P_3 \approx 1.3893346/1.0266331946^{101}$. The type I error rate is then approximately equal to $P_3 - P_2 \approx 0.00521934$. For comparison, a 99% confidence interval obtained by performing 10 billion random simulations is $0.0052166 - 0.0052203$.

*

Figure 25 illustrates the precision of this estimate for different values of the substitution rate κ and of the read size k . The approximation is grossly inaccurate for $\kappa = 0.05$, where it is even negative for reads under 40 nucleotides. The issue here is slow convergence, as mentioned in section 3.7. The dominant singularity is approximately equal to 1.066122, while the modulus of the second and the third is approximately equal to 1.110296. For $k \leq 40$, the contribution of the higher order singularities is not negligible; using only the dominant singularity we obtain $P_3 \approx 0.158993$, compared to $P_3 \approx 0.163655$ when using three singularities.

Surprisingly, the estimations of P_3 in the cases $\kappa = 0.15$ and $\kappa = 0.25$ are equally inaccurate, but the errors almost perfectly compensate with those on the approximation of P_2 . Even more surprisingly, using three singularities improves the approximations of type I error rates for $\kappa = 0.05$, but makes them worse for $\kappa = 0.15$ and $\kappa = 0.25$ because the errors on P_2 and P_3 no longer compensate. An empirical solution is to record the (p, κ) pairs for which to use one or three singularities. Even though this solution is not satisfactory from the theoretical point of view, it gives accurate approximations for the whole range of parameters, as shown in figure 26.

We cannot use the method presented in section 4.2 to extend the model to more than one duplicate because conditional independence does not hold for MEM seeds. But we can still use the estimates above as a lower bound for the probability of type I error. If the proportion of the genome that is repeated is x and P^* is the maximum of $P_3 - P_2$ relative to unknown κ , then xP^* is again a reasonable lower bound on the probability of type I errors.

5 Average quantities

5.1 General approach

The analytic combinatorics approach allows computing the average of many quantities of interest. In the seeding problem, one such quantity is the number of errors for reads of different kinds. So far, weighted generating functions only

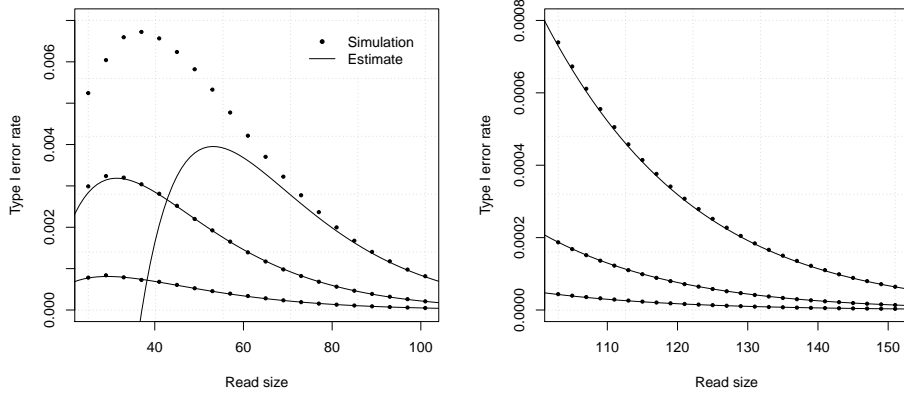


Figure 25: **Example estimates of MEM seeding probability (one duplicate).** The analytic combinatorics estimates are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size will give a type I error when using MEM seeds, either estimated by 10,000,000 (left) or 100,000,000 (right) random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$ and $p = 0.05$, $\kappa = 0.05$, $\kappa = 0.15$ or $\kappa = 0.25$ (from top to bottom).

marked the size of the reads through the variable z . We now need to introduce another variable to mark the second quantity, which means that we will deal with bivariate weighted generating functions. For concreteness, if $a_{k,n}$ is the total weight of seedless reads of size k and with n errors, the average number of errors for reads of size k is by definition

$$\left(\sum_{n=0}^{\infty} n a_{k,n} \right) / \left(\sum_{n=0}^{\infty} a_{k,n} \right). \quad (27)$$

To compute this quantity, we introduce the variable u marking the number of substitutions and we write

$$W(z, u) = \sum_{k=0}^{\infty} \sum_{n=0}^{\infty} a_{k,n} z^k u^n.$$

Finding an explicit formula for $W(z, u)$ is the focus of sections 5.2 to 5.4. For now, observe that (27) can be expressed from $W(z, u)$ as follows. On the one hand, we have

$$W(z, 1) = \sum_{k=0}^{\infty} \left(\sum_{n=0}^{\infty} a_{k,n} \right) z^k.$$

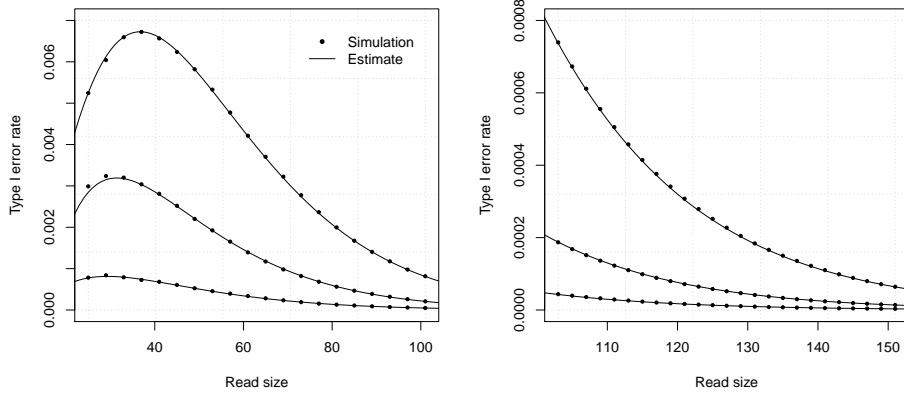


Figure 26: **Example ad hoc estimates of MEM seeding probability (one duplicate).** The analytic combinatorics estimates with *ad hoc* adjustments are benchmarked against random simulations. Shown on both panels are the probabilities that a read of given size will give a type I error when using MEM seeds, either estimated by 10,000,000 (left) or 100,000,000 (right) random simulations (dots), or by the method described above (lines). The curves are drawn for $\gamma = 17$ and $p = 0.05$, $\kappa = 0.05$, $\kappa = 0.15$ or $\kappa = 0.25$ (from top to bottom). The simulation data points are the same as those shown in figure 25. Three singularities are used to approximate P_3 for $\kappa = 0.05$, and only one for $\kappa = 0.15$ and $\kappa = 0.25$.

So the denominator of (27) is the coefficient of z^k in $W(z, 1)$. On the other hand, by taking the derivative of $W(z, u)$ with respect to u , and setting $u = 1$, we obtain

$$W_u(z, 1) = \frac{\partial W(z, u)}{\partial u} \Big|_{u=1} = \sum_{k=0}^{\infty} \left(\sum_{n=0}^{\infty} n a_{k,n} \right) z^k.$$

So the numerator of (27) is the coefficient of z^k in $W_u(z, 1)$. Note that $W(z, 1)$ and $W_u(z, 1)$ are actually univariate weighted generating functions, so we can use the methods developed earlier to obtain asymptotic estimates for their coefficients. However, proposition 5 will not apply to approximate the coefficients of $W_u(z, 1)$. Since all the weighted generating functions considered here are ratios of polynomials of the form $W(z, u) = P(z, u)/Q(z, u)$, the derivative $W_u(z, 1)$ can be expressed as $P_u(z, 1)/Q(z, 1) - P(z, 1)Q_u(z, 1)/Q(z, 1)^2$. Because of the term $Q(z, 1)^2$ at the denominator, the singularities of $W_u(z, 1)$ are not simple poles.

Proposition 10 below shows how to extract the coefficients of weighted generating functions in this particular case.

Proposition 10. *If a function $W(z)$ is the ratio of two polynomials $P(z)/Q(z)^2$, and Q has only simple roots with distinct moduli, then*

$$[z^k]W(z) \sim \left((k+1) \frac{P(z_1)}{z_1 Q'(z_1)^2} - \frac{P'(z_1)}{Q'(z_1)^2} + \frac{P(z_1)Q''(z_1)}{Q'(z_1)^3} \right) \frac{1}{z_1^{k+1}}, \quad (28)$$

where z_1 is the root of Q with smallest modulus.

As in proposition 5, we start by proving the lemma that will give the functional expression of the asymptotic expansion.

Lemma 3. For $|z| < a$ we have

$$\frac{1}{(1-z/a)^2} = \sum_{k=0}^{\infty} (k+1) \frac{z^k}{a^k}. \quad (29)$$

Proof.

$$\frac{1}{(1-z/a)^2} = a \frac{\partial}{\partial z} \left(\frac{1}{1-z/a} \right) = a \sum_{k=0}^{\infty} \frac{kz^{k-1}}{a^k},$$

where the last equality is obtained by differentiating after applying lemma 2. \square

We now prove proposition 10.

Proof. As in the proof of proposition 5, let z_1, z_2, \dots, z_n be the complex roots of Q , sorted by increasing order of *modulus*. Since the roots of $Q(z)^2$ all have multiplicity 2, there exists constants $\alpha_1, \dots, \alpha_n$ and β_1, \dots, β_n such that the partial fraction decomposition of the rational function $P(z)/Q(z)^2$ can be written as

$$\frac{P(z)}{Q(z)^2} = \sum_{j=1}^n \frac{\alpha_j}{(z-z_j)^2} + \frac{\beta_j}{z-z_j} = \sum_{j=1}^n \frac{\alpha_j/z_j^2}{(1-z/z_j)^2} - \frac{\beta_j/z_j}{1-z/z_j}.$$

As in the proof of proposition 5, we assumed without loss of generality that the degree of P is lower than the degree of Q^2 . Now applying lemmas 2 and 3, we see that the coefficient of z^k in $P(z)/Q(z)^2$ can be expressed as

$$\sum_{j=1}^n (k+1) \frac{\alpha_j}{z_j^{k+2}} - \frac{\beta_j}{z_j^{k+1}}. \quad (30)$$

The sum above is dominated by the term with the highest exponential rate of increase, *i.e.* $j = 1$ because z_1 has smallest modulus by definition. Thus, the coefficient of z^k in $P(z)/Q(z)^2$ is asymptotically equivalent to

$$(k+1) \frac{\alpha_1}{z_1^{k+2}} - \frac{\beta_1}{z_1^{k+1}}.$$

We now need to find the values of α_1 and β_1 . As z_1 is a root of Q , there exists a polynomial $Q_1(z)$ such that

$$Q(z) = (z - z_1)Q_1(z). \quad (31)$$

We can thus write $P(z)/Q(z)^2$ as

$$\frac{P(z)}{(z - z_1)^2 Q_1(z)^2} = \frac{\alpha_1}{(z - z_1)^2} + \frac{\beta_1}{z - z_1} + \sum_{j=2}^n \frac{\alpha_j}{(z - z_j)^2} + \frac{\beta_j}{z - z_j}. \quad (32)$$

Multiplying both sides of (32) by $(z - z_1)^2$ and setting $z = z_1$ shows that $\alpha_1 = P(z_1)/Q_1(z_1)^2$. To find the value of $Q_1(z_1)$ we differentiate (31) and let $z = z_1$, yielding $Q'(z_1) = Q_1(z_1)$. We thus obtain $\alpha_1 = P(z_1)/Q'(z_1)^2$.

To find the value of β_1 , we subtract $\alpha_1/(z - z_1)^2$ on both sides of (32) and obtain

$$\frac{P(z) - P(z_1)Q_1(z)^2/Q_1(z_1)^2}{(z - z_1)^2 Q_1(z)^2} = \frac{\beta_1}{z - z_1} + \sum_{j=2}^n \frac{\alpha_j}{(z - z_j)^2} + \frac{\beta_j}{z - z_j}. \quad (33)$$

Since $P(z_1) - P(z_1)Q_1(z_1)^2/Q_1(z_1)^2 = 0$, there exists a polynomial $Q_2(z)$ such that

$$P(z) - P(z_1)Q_1(z)^2/Q_1(z_1)^2 = (z - z_1)Q_2(z). \quad (34)$$

Multiplying (33) by $z - z_1$ and setting $z = z_1$, we obtain $\beta_1 = Q_2(z_1)/Q_1(z_1)^2$. To find the value of $Q_2(z_1)$ we differentiate (31) two times and (34) one time, finally obtaining $\beta_1 = P'(z_1)/Q'(z_1)^2 - P(z_1)Q''(z_1)/Q'(z_1)^3$, which concludes the proof. \square

Remark 21. Expression (28) is asymptotically equivalent to the simpler expression

$$(k+1) \frac{P(z_1)}{Q'(z_1)^2} \frac{1}{z_1^{k+2}}. \quad (35)$$

However, the convergence of expression (35) is slow. Indeed, dividing the exact expression (30) by (35) gives an error term in $O(1/k)$. In comparison, dividing (30) by (28) gives an error term in $O(|z_1/z_2|^k)$, which decreases exponentially, as in proposition 5.

We are now ready to approximate the mean number of errors in reads of different kinds. We treat the distinct types of errors separately, in the framework of the corresponding error model from section 3.3 to 3.5.

5.2 Substitutions

Applying a now familiar strategy, we write the generating function of simple objects that we combine them into more complex objects. Recall from section 3.3 that in the uniform substitution model, the weighted generating function of substitutions is pz , where p is the error rate. In order to mark them with the variable u , we replace pz by pzu . For every substitution, the power of z increases

by 1, so the size of the read increases by 1. The power of u also increases by 1 irrespective of the size, which allows us to mark the number of substitutions.

We could derive the weighted generating function by the transfer matrix method described in section 3.3, but we can get an immediate result by using equation (12), where we observed that the weighted generating function of reads without exact seed can be expressed as

$$\frac{1 + F_\gamma(z)}{1 - pz(1 + F_\gamma(z))}. \quad (12)$$

Here the weighted generating function of substitutions appears explicitly as pz . We can replace it by pzu to obtain directly

$$S(z, u) = \frac{1 + F_\gamma(z)}{1 - pzu(1 + F_\gamma(z))}. \quad (36)$$

$S(z, 1)$ is simply the weighted generating function of seedless reads derived in section 3.3, and for which we already know the coefficient asymptotics. So we already have the denominator of (27). Now differentiating (36) with respect to u , we obtain

$$\left. \frac{\partial}{\partial u} \left(\frac{1 + F_\gamma(z)}{1 - pzu(1 + F_\gamma(z))} \right) \right|_{u=1} = \frac{pz(1 + F_\gamma(z))^2}{(1 - pz(1 + F_\gamma(z)))^2}. \quad (37)$$

As mentioned above, the form of the weighted generating function calls for using proposition 10 (and not proposition 5). The dominant singularity is the same for both the numerator and the denominator of (27), so the terms in z_1^{k+1} cancel out. What remains is an expression of the form $C_1 k + C_2$, where C_1 and C_2 are constants. In other words, the average number of substitutions in reads without exact seed increases as an affine function of the size. We make this result more accurate in the following proposition.

Proposition 11. *Under the assumptions of the error model of section 3.3, the average number of substitutions in seedless reads of size k is asymptotically equivalent to*

$$\frac{(1 - qz)(C_1(k + 1) - C_2)}{1 - (\gamma + 1 - \gamma qz_1)(qz_1)^\gamma},$$

with

$$C_1 = 1 - (qz_1)^\gamma, \text{ and}$$

$$C_2 = \frac{(1 - (2 - (1 - qz_1)\gamma^2 + 2\gamma)(qz_1)^\gamma + (1 + (1 - qz_1)\gamma^2 - 2\gamma)(qz_1)^{2\gamma})}{1 - (\gamma + 1 - \gamma qz_1)(qz_1)^\gamma},$$

and where z_1 is the only positive root of the polynomial $1 - pz(1 + F_\gamma(z))$.

Proof. Apply proposition 10 to (37), use proposition 7 and simplify. \square

Remark 22. The estimate obtained from (35) instead of (28) increases linearly with k . In this case, only the relative error vanishes asymptotically. The absolute error remains constant.

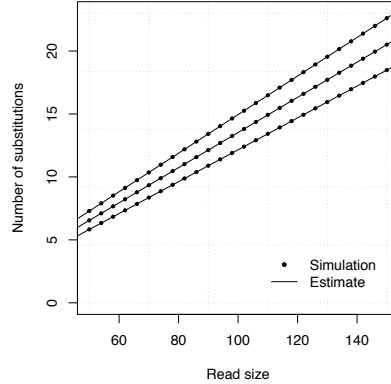


Figure 27: **Estimating the average number of substitutions.** The average number of substitutions in reads without γ -exact seed is shown for different read sizes, either estimated by 10,000,000 random simulations (dots), or by analytic combinatorics (lines). The curves are drawn for $\gamma = 17$, $p = 0.08$, $p = 0.10$ or $p = 0.12$ (from bottom to top).

Figure 27 illustrates the accuracy of the estimate. The affine relationship between the average number of errors and the size of the read without exact seed is apparent, and the approximations are very close to the target values.

Remark 23. What about the average number of substitutions in reads that contain a γ -exact seed? We can compute it indirectly from the quantities that we already have defined.

The average number of substitutions for reads of size k (regardless of whether they contain a seed) is pk . Call P_1 the probability that a read of size k contains no seed and E the average number of substitutions in such reads. The quantity of interest, x follows the relationship

$$P_1E + (1 - P_1)x = pk.$$

From this we see that the average number of substitutions in reads that contain a γ -exact seed is $(pk - P_1E)/(1 - P_1)$. This relationship is not affine because P_1 decreases exponentially as k increases, but for long reads P_1 vanishes so it is approximately linear and equal to pk .

5.3 Deletions

To count the average number of deletions, we take the weighted generating function of the reads without seed in the error model of section 3.4, namely

$$\frac{1 + (1 - \delta)F_\gamma(z)}{1 - pz - (pz(1 - \delta) + \delta)F_\gamma(z)}. \quad (17)$$

Here it is important to remember that for simplicity, equation (17) ignores the deletions that occur immediately before and after substitutions, because they have no impact on the presence of a seed. For the purpose of counting deletions, we need to take them all into consideration, but this requires working with the proper weighted generating function.

A read can be thought of as a walk on the graph shown in figure 28. Even though deletions appear as an explicit state, their size is 0 and they occur only between nucleotides (which implies that a read can neither start nor end with a deletion). Also, a deletion must be followed by either a match or a substitution; it cannot be followed by a deletion. The reason is that a deletion has size 0 regardless the number of deleted nucleotides. In the symbolic representation of the read, either there is a deletion of any size between two nucleotides, or there is no deletion.

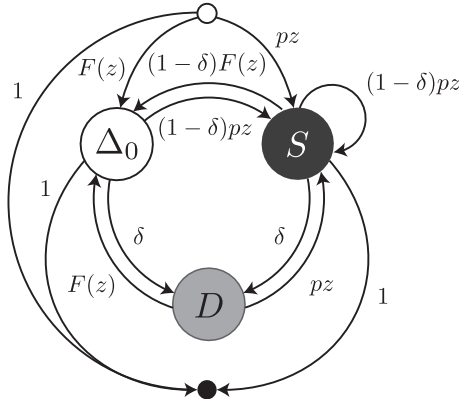


Figure 28: **Transfer graph with explicit deletions.** Reads are viewed as sequences of error-free intervals (symbol Δ_0) or substitutions (symbol S) or deletions (symbol D), with respective weighted generating functions $F(z)$, pz and δ .

Here, reads are sequences of error-free intervals with symbol Δ_0 and weighted generating function $F(z)$ or substitutions with symbol S and weighted generating function pz , or deletions with symbol D and weighted generating function δ . The only forbidden transitions in the sequence are that two error-free intervals cannot follow each other (together they form a single error-free interval) and two deletions cannot follow each other (together they form a single deletion). In contrast with figure 8 where deletions do not have their own vertex, we need to

account for the absence of deletions in transitions between error-free intervals and substitutions. This is done by prepending a weight equal to $(1 - \delta)$.

As in the previous section, in order to count deletions with parameter v we weight their generating function by δv . For every deletion the power of v increases by 1 (and the power of z remains the same). This allows us to mark the number of deletions. The transfer matrix then becomes a function of z , marking the size of the reads, and of v , marking the number of deletions. The final expression of the transfer matrix is

$$M(z, v) = \begin{matrix} & \Delta_0 & S & D \\ \Delta_0 & 0 & (1 - \delta)pz & \delta v \\ S & (1 - \delta)F(z) & (1 - \delta)pz & \delta v \\ D & F(z) & pz & 0 \end{matrix}.$$

Recall that reads can neither start nor end by a deletion because they are undetectable at those positions. Thus the head vector is $H(z, v) = (F(z), pz, 0)^\top$ and the tail vector is $T(z, v) = (1, 1, 0)^\top$. The only sequence we need to add is the empty sequence ε , so $\psi(z, v) = 1$. Following proposition 4, the weighted generating function $S(z, v)$ is equal to $\psi(z, v) + H(z, v)^\top \cdot (I - M(z, v))^{-1} \cdot T(z, v)$.

Following the general strategy to count average quantities, we first compute $S(z, 1)$. Naturally, we obtain expression (17), which is the weighted generating function of reads without exact seed. We also differentiate $S(z, v)$ with respect to v and let $v = 1$ to obtain

$$S_v(z, 1) = \frac{\delta(pz(1 + (1 - \delta)F_\gamma(z)) + F_\gamma(z))^2}{(1 - pz - (pz(1 - \delta) + \delta)F_\gamma(z))^2}. \quad (38)$$

As in the section 5.2, we derive the asymptotics by applying proposition 10 to (38) and by applying proposition 5 to (17). The terms z_1^{k+1} cancel out when computing the ratio as per equation (27), so the number of deletions in reads without seed is asymptotically affine, *i.e.* of the form $C_1 k + C_2$, where C_1 and C_2 are constants. In both cases, C_1 and C_2 have cumbersome expressions, but they are easy to compute from propositions 5 and 10.

The accuracy of the estimates is illustrated in figure 29. Once again the affine relationship is a good fit and the approximations are close the target values.

5.4 Insertions

Finally, we compute the average amount of insertions for reads without exact seed in the full error model of section 3.5. The simplest way to do this is to mark insertions directly in the transfer matrix where they appear explicitly. For this, we append the extra variable w to their weighted generating functions and obtain the following transfer matrix

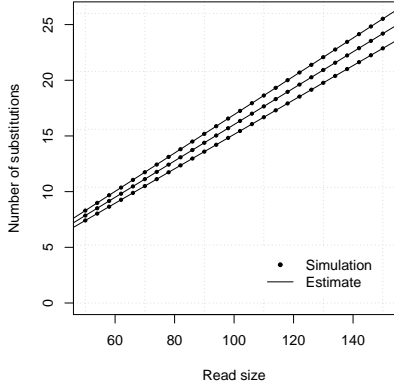


Figure 29: **Example estimates of average number of deletions.** The average number of deletions in reads without γ -exact seed is shown for different read sizes, either estimated by 10,000,000 random simulations (dots), or by analytic combinatorics (lines). The curves are drawn for $\gamma = 17$, $p = 0.05$ and $\delta = 0.14$, $\delta = 0.15$ or $\delta = 0.16$ (from bottom to top).

$$M(z, w) = \begin{matrix} \Delta_0 & S & I \\ \Delta_0 & \delta F_\gamma(z) & pz & rzw \\ S & F_\gamma(z) & pz & rzw \\ I & \frac{1-\tilde{r}}{1-r} F_\gamma(z) & \frac{1-\tilde{r}}{1-r} pz & \tilde{r} zw \end{matrix}.$$

In the expression above, we have used $F_\gamma(z)$ instead of $F(z)$ because we are directly considering reads without seed. We also update the head and tail vectors $H(z, w) = (F_\gamma(z), pz, rzw)^\top$ and $T(z, w) = (1, 1, 1)^\top$, and we add the empty sequence, which does not contain any insertion, so $\psi(z, w) = 1$. Following proposition 4, $S(z, w) = \psi(z, w) + H(z, w)^\top \cdot (I - M(z, w))^{-1} \cdot T(z, w)$.

As explained in section 5.1, we obtain the average number of insertions by computing ratio (27) from the coefficients of $S(z, 1)$ and of $S_w(z, 1)$. Computing $S(z, 1)$ yields expression (21), *i.e.* the weighted generating function of reads without exact seed. By differentiating $S(z, w)$ and setting $w = 1$ we find $S_w(z, 1)$ to be equal to

$$\frac{rz(1-r)(1+(1-\delta)F(z))(1-r-pz(\tilde{r}-r)+c(z)F(z))}{(1-a(z)-b(z)F_\gamma(z))^2}, \quad (39)$$

where $a(z)$ and $b(z)$ are defined as in (20), and where $c(z)$ is a first degree polynomial defined as

$$c(z) = 1 - \tilde{r} - \delta(1-r) - (1-\delta)(\tilde{r}-r)pz.$$

As in the section 5.2, we derive the asymptotics by applying proposition 10 to (39) and by applying proposition 5 to (21). The terms z_1^{k+1} cancel out when computing the ratio as per equation (27), so the number of insertions in reads without seed is asymptotically affine, *i.e.* of the form $C_1 k + C_2$, where C_1 and C_2 are constants. In both cases, C_1 and C_2 have cumbersome expressions, but they are easy to compute from propositions 5 and 10.

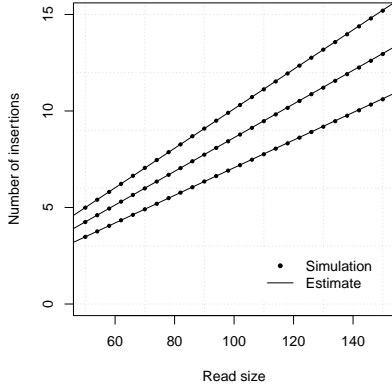


Figure 30: **Example estimates of average number of insertions.** The average number of insertions in reads without γ -exact seed is shown for different read sizes, either estimated by 10,000,000 random simulations (dots), or by analytic combinatorics (lines). The curves are drawn for $\gamma = 17$, $p = 0.05$, $\delta = 0.15$, $\tilde{r} = 0.45$ and $r = 0.04$, $r = 0.05$ or $r = 0.06$ (from bottom to top).

The accuracy of the estimates is illustrated in figure 30. Once again the affine relationship is a good fit and the approximations are close the target values.

It is also meaningful to compute the average number of substitutions or deletions in the full error model. For this, one can start from the transfer matrix, mark the quantity of interest and proceed with the general strategy exposed in section 5.1.

To compute the average number of substitutions, we use the transfer matrix

$$M(z, u) = \begin{matrix} & \Delta_0 & S & I \\ \Delta_0 & \left[\begin{array}{ccc} \delta F_\gamma(z) & pzu & rz \\ F_\gamma(z) & pzu & rz \\ \frac{1-\tilde{r}}{1-r} F_\gamma(z) & \frac{1-\tilde{r}}{1-r} pzu & \tilde{r}z \end{array} \right] \end{matrix}.$$

To compute the average number of deletions, we use the transfer matrix

$$M(z, v) = \begin{matrix} & \Delta_0 & S & D & I \\ \Delta_0 & 0 & (1-\delta)pz & \delta v & rz \\ S & (1-\delta)F_\gamma(z) & (1-\delta)pz & \delta v & rz \\ D & F_\gamma(z) & pz & 0 & 0 \\ I & \frac{1-\tilde{r}}{1-r}F_\gamma(z) & \frac{1-\tilde{r}}{1-r}pz & 0 & \tilde{r}z \end{matrix}.$$

Finally, we can also consider all errors together (deletions always counting as a single error). For this we mark all the types of errors with the same variable, say w . In this case, we use the transfer matrix

$$M(z, w) = \begin{matrix} & \Delta_0 & S & D & I \\ \Delta_0 & 0 & (1-\delta)pzw & \delta w & rzw \\ S & (1-\delta)F_\gamma(z) & (1-\delta)pzw & \delta vw & rzw \\ D & F_\gamma(z) & pzw & 0 & 0 \\ I & \frac{1-\tilde{r}}{1-r}F_\gamma(z) & \frac{1-\tilde{r}}{1-r}pzw & 0 & \tilde{r}zw \end{matrix}.$$

In all the cases, we obtain an affine relationship between the size of the read and the average number of error of a given type.

6 Conclusion

This concludes our introductory tour of the applications of analytic combinatorics to seeding methods. There is of course much more to say. Our purpose here is to show how the general strategy of analytic combinatorics gives reasonable solutions to problems that were previously difficult to address.

When using exact seeds, inexact seeds or MEM seeds, the estimates of the seeding probabilities are robust and relatively straightforward. The data collected during the alignment can be used to estimate the parameters of the error model to auto-tune the seeding heuristic during the run. This gives tight control of the type II error rate. More challenging is to control the type I error rate. The estimates show some numeric instability when using MEM seeds, but the real difficulty is the lack of theoretical background on the seeding process when the target is duplicated. Significant progress on this line may not come from analytic combinatorics, but from more appropriate data structures or algorithms to seed such sequences more efficiently.

Seeding is not only used in mapping, but also in other alignment problems. In this regard, the work presented above can be applied to different contexts. That said, mapping high throughput sequencing reads is a “sweet spot” for analytic combinatorics because the sequences are long enough for the approximations to be accurate.

Finally, the concepts developed above can also be used to sample reads of different kinds. The analogy between reads and walks on graphs with weighted edges allows us to generate reads by simulating a random walk. The trick is to

convert the transfer matrix into a transition matrix by using the weights of the objects as their probabilities of occurrence.

In summary, analytic combinatorics is a powerful strategy that comes with a rich toolbox with many applications in modern bioinformatics. We expect more applications to see the light as new algorithms and new heuristics are developed in the future.

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