ABSTRACT

Motivation: Currently, re-sequencing approaches use multiple modules serially to interpret raw sequencing data from next-generation sequencing platforms, while remaining oblivious to the genomic information until the final alignment step. Such approaches fail to exploit the full information from both raw sequencing data and the reference genome that can yield better quality sequence reads, SNP-calls, variant detection, as well as an alignment at the best possible location in the reference genome. Thus, there is a need for novel reference-guided bioinformatics algorithms for interpreting analog signals representing sequences of the bases \((A, C, G, T)\), while simultaneously aligning possible sequence reads to a source reference genome whenever available.

Results: Here, we propose a new base-calling algorithm, \textsc{TotalRecaller}, to achieve improved performance. A linear error model for the raw intensity data and Burrows–Wheeler transform (BWT) based alignment are combined utilizing a Bayesian score function, which is then globally optimized over all possible genomic locations using an efficient branch-and-bound approach. The algorithm has been implemented in soft- and hardware [field-programmable gate array (FPGA)] to achieve real-time performance. Empirical results on real high-throughput Illumina data were used to evaluate \textsc{TotalRecaller}'s performance relative to its peers—Bustard, BayesCall, Ibis and Rolexa—based on several criteria, particularly those important in clinical and scientific applications. Namely, it was evaluated for (i) its base-calling speed and throughput, (ii) its read accuracy and (iii) its specificity and sensitivity in variant calling.

Availability: A software implementation of \textsc{TotalRecaller} as well as additional information, is available at: http://bioinformatics.nyu.edu/wordpress/projects/totalrecaller/

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1 INTRODUCTION

Recent advances in sequencing technology continue to introduce novel and diverse high-throughput DNA sequencing platforms, such as Illumina. These innovations, promise to revolutionize biological, biomedical and translational research by rapidly transferring genomics breakthroughs from the byte banks to bedside. For any large-scale genomic sequencing effort aiming to fulfill these goals, an important and rate-limiting initial step involves Base-Calling, which is now assuming an even more critical role, especially for these new (next- and next-next-generation) sequencing platforms. When the source genome references are available, the sequence-reads obtained by the base-caller is next subjected to alignment (followed by variant-caller) procedures successively—although, in principle, all three steps could be carried out in one integrated step while promising improvements in accuracy, computational complexity, data storage and transmission.

Base-calling takes the vector analog time series of signals generated by the sequencing machines as input, and produces a base-by-base digitized estimate of the underlying DNA sequence that is most likely to have given rise to those signals. Although the next-generation sequencing technologies listed above have reduced the cost and increased the throughput, these platforms pose new challenges for base-calling, as their technology is based on either relatively small number of unsynchronized molecules (e.g. Illumina, 454 Life Science, Ion Torrent, SOLiD) or single molecules with weak signal (e.g. Pacific Biosciences, Oxford Nanopore, Helicos, Life Technologies). As a consequence, these platforms are not only error prone, they also corrupt signals in the data by non-stationary errors and generate much shorter reads than what is needed for both proper alignment and sequence assembly, as well as what used to be routinely possible with the traditional Sanger sequencers.

Motivated by such challenges, novel base-calling frameworks have been proposed to deal with many unknown sources of noise in these data. These methods have already demonstrated that considerable improvement both in quality and read length is possible through sophisticated signal processing methods such as statistical learning (Kao et al., 2009), supervised learning and support vector machine (SVM) (Erlich et al., 2008; Kircher et al., 2009), and model-based clustering and information theory (Rougemon et al., 2008).

However, in doing so, these methods also expose Base-Calling to several other limitations: (i) Over-fitting: parametric models (e.g. Alta-Cyclic, Ibis, BayesCall, Rolexa) are likely to suffer from over-fitting to the in-sample data and thus are unlikely to be very robust in dealing with varying out-of-sample datasets, even from the same sequencing platforms. (ii) Computational cost: in a preprocessing step, all base-callers must learn the error model in the training data in order to build a classifier that then corrects the errors in the signal. This preprocessing step can be very time consuming (as in the case of BayesCall, Alta-Cyclic and Ibis) and may require a cluster computer facility (as in the case of Alta-Cyclic), thus preventing them from real-time base-calling, as would be needed.
in many clinical applications. (iii) Training phase: since a subclass of these base-callers (e.g. Alta-Cyclic, Ibis) needs a training phase using a library of ‘correct’ reads, they require both a secondary base-caller and a sequence aligner to get started with the training library. On the contrary, this concern has been somewhat alleviated in the newer base-callers (e.g. Bustard, BayesCall), which estimate their parameters solely from intensity files. (iv) Technology dependent: many base-callers (e.g. Alta-Cyclic, Ibis and BayesCall) use a detailed parametric model to describe the signal distortion as a function of successive cycles. Such models require and hard-wire specific knowledge of the underlying sequencing technology into the algorithm, thus making it harder to customize the base-caller to support other platforms.

Currently, a major application of the next-generation sequencing technologies is in re-sequencing (e.g. DNASeq: DNA sequencing with known reference genome; RNASeq: mRNA sequencing with known gene isoforms; CHIPSeq: chromatin immunoprecipitation sequencing with known binding sites). Despite the obvious centrality of alignment in these applications, traditional base-callers have avoided performing alignment until the end of the base-calling process; in fact, the typical pipeline for a re-sequencing process traditionally consists of two sequential steps:

1. **Base-calling**: each single base of the read is called according to the intensity signal and error profiles.

2. **Alignment**: sequence reads are aligned to a reference genome.

Because the base-calling process is error prone, and because correct alignment to the reference genome is non-trivial, high coverage is required in order to reduce the errors in re-sequencing and recover the true full DNA sequence.

To perform the alignment to the reference genome, the standard Smith–Waterman was developed in the context of homology analysis, is based on dynamic programming and thus requires its score function to evaluate the alignment quality to satisfy ‘the principle of optimality’ (Bellman et al., 1959; Dreyfus, 2002). As the Illumina sequencing pipeline has been described elsewhere (Carrington et al., 2007; Metzker, 2005; Bentley, 2006), we report here, for convenience, only a minimal description—essentially what is needed in this article to understand the source of errors in the data and how it is handled by TOTALReCaller. The Illumina pipeline, generating raw signals prior to base-calling and alignment, consists of five steps in which: (i) DNA sample is prepared; (ii) pieces of DNA in the sample are randomly fragmented and placed on a flow cell; (iii) fragments are amplified into clusters. Each cluster consists out of ~1000 identical strands; (iv) fluorescent markers, lasers and CCD sensors are used to read the clusters base by base (so called cycles), resulting in four images for each cycle; and finally (v) the images are analyzed and a single analog intensity value is determined for each image. Base-calling is the process of determining the correct sequence of bases \( b \in \{A, C, G, T\}^T \) from a sequence of images, one for each base \( B \in \Sigma = \{A, C, G, T\} \). While there are many challenges faced by the base-callers attempting to correctly interpret Illumina intensities data, the major one is due to relatively low signal-to-noise ratio (SNR) of the output, which worsens precipitously with the cycle numbers—(ascribed primarily to polymerase desynchronization and its low processivity. Figure 1...
highlights the magnitude of this problem by showing the intensity for each channel averaged over a large number (~5 x 10^5) of 78 bp sequence reads for the bacteriophage phiX genome. As shown in the figure, calling the bases A and G from their intensity channels does not pose a severe problem, at least for the first 40 cycles. In contrast, channels C and T appear hopelessly disrupted within the first few cycles.

Source of errors in Illumina raw sequencing data: as reported previously in Erlich et al. (2008), there are four dominant sources of noise affecting the intensities generated by Illumina:

1. Crosstalk: the intensity channels are not independent. This inter-dependence is due to the fact that the fluorescent markers for A, C and G, T emit photons with similar wavelengths and get excited by the same lasers, and because fluorescent markers from one cycle can only be chemically partially removed (“washed”) before the cycles for the next nucleotides (all performed in the same flow cell).

2. Fading: with subsequent cycles, the absolute intensity of light emitted from the cluster of DNA strands diminishes because fluorescent markers are only able to bind to fewer and fewer strands within the clusters.

3. Lagging (Phasing): some strands in the clusters start to lag behind the population, as in each cycle some of the polymerases fail to operate synchronously, but then rejoin the other strands in subsequent cycles, whence producing ambiguous intensity values. Eventually, the correct channel gets obscured by the other wrong channel intensities, leading to wrong base-calls.

4. Leading (Pre-phasing): some strands in the clusters start to lead ahead of the population, which also causes ambiguous and incorrect channel intensity values in a fashion analogous to lagging.

These noise factors dominate and affect the signal differently in different cycles: in the first few cycles, cross-talk is the major source of base-call errors; however, in later cycles, fading, lagging and leading prevail. We have observed that lagging often causes many false-positive insertions in the distal extending-end of sequence reads: in later cycles, intensities measured in cycle k more and more reflect what would have been the value in cycle (k–1). This process leads directly to ‘base-stuttering’, which occurs much more frequently after some threshold value for k, the cycle number. This dynamic can be modeled by a step function appearing randomly but more frequently in later cycles, thus making it extremely difficult to analyze. This effect has important implications for the succeeding alignment step, since many popular short-read sequence aligners cannot align gapped sequence reads [Langmead et al. (2009); Li and Durbin (2009)]. We have observed that in our datasets the effects of leading on the SNR are negligible in comparison to the other three causes of error (cross-talk, fading and lagging).

2 RESULTS

Including the base-caller introduced in this article, currently there are six base-callers for Illumina sequencing machines: TOTALReCaller, Bustard (Illumina), Alta-Cyclic [CSHL, Erlich et al. (2008)], BayesCall [UC Berkeley, Kao et al. (2009)], Ibis [Max Planck, Kircher et al. (2009)] and Rolexa [Université de Lausanne, Rougemont et al. (2008)]. In the following, we compare the performance of these base-callers according to the same metrics that have been used in the literature: namely, base-calling error rate, alignment rate and base-calling speed. Since TOTALReCaller utilizes a reference genome for base-calling, base-calling statistics from datasets of three different organisms (with reference genomes of varying qualities) were analyzed (Table 1). Furthermore, base-calling results of TOTALReCaller are presented with different weights on reference alignment with respect to intensity information. Finally, we conclude this section with detailed statistics on TOTALReCaller’s SNP sensitivity and specificity, as they would be the most important in many clinical applications.

The base-calling error rate: This measures the quality per cycle (base pair position) of the sequence reads produced by a given base-caller (Fig. 2). In order to generate error rates based on the same set of reads for each of the base-callers, we aligned all Bustard reads to the respective reference genome (Table 1) in order to create a set of ‘correct reads’. We then perform a base-by-base comparison between this set of ‘correct reads’ and the sequence reads created by each of the base-callers, resulting in an error rate for each position (cycle) in a sequence read. Since we used the output of Bustard to create the set of correct reads, we introduced a bias, favoring Bustard. For the small genomes, phiX and E.coli, we used the aligner BLAT (Kent, 2002), which allows accurate, gapped alignment to create the set of ‘correct reads’. For the poplar dataset, we used Bowtie (Langmead et al., 2009) to create the set of ‘correct reads’. We had to use Bowtie instead of BLAT in order to properly handle the current draft of the poplar genome (Tuskan et al., 2006) genome, which is of relatively lower quality in comparison to E.coli and phiX, e.g. poplar consists of many contigs (out of 2518) that have not been yet phased to a scaffold (http://www.phytozone.net/poplar). The low quality, in conjunction with the length and complex structure of the poplar genome, results in an unusually large number of false positive alignments, which, when analyzed by a sensitive aligner, makes it

Table 1. Datasets

<table>
<thead>
<tr>
<th>Genome</th>
<th>Genome size</th>
<th>No. of clusters</th>
<th>No. of cycles</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>phiX</td>
<td>~5.4kB</td>
<td>11803171</td>
<td>78</td>
<td>One lane of Genome Analyzer I run.</td>
</tr>
<tr>
<td>E.coli</td>
<td>~4.5MB</td>
<td>35027442</td>
<td>125</td>
<td>The second pair of a paired-end Genome Analyzer II run (one lane).</td>
</tr>
<tr>
<td>poplar</td>
<td>~420MB</td>
<td>31445866</td>
<td>109</td>
<td>The first pair of a paired-end Genome Analyzer II run (one lane).</td>
</tr>
</tbody>
</table>

Since TOTALReCaller injects knowledge from a reference genome into the base-calling process, base-calling becomes dependent on the genome structure. Therefore, we evaluated TOTALReCaller's performance on three different datasets: phiX, E.coli and poplar.
impossible to create a valid set of ‘correct reads’. Since Bowtie and other suffix tree-based algorithms are generally less sensitive than BLAST-like (Altschul et al., 1997) alignment algorithms (e.g. they do not allow gapped alignment), they produce fewer but significantly more accurate sets of ‘correct reads’, especially in the case of a ‘bad’ reference genome. The base-calling error rates based on the reads produced by the base-callers and the set of ‘correct reads’ can be found in Figure 2.

It is shown that base sequence reads generated by TotalReCaller have a significantly lower base-calling error rate in comparison to all other base-callers. Furthermore, the error rate can be controlled by choosing an alignment weight $w_{\text{align}}$. However, it is necessary to understand that increasing the alignment weight $w_{\text{align}}$ does have negative influences which are discussed together with SNP sensitivity and specificity. Based on the previous discussion, it is safe to conclude that the error rates for the poplar dataset may be used only for a qualitative (and not a quantitative) comparison.

**The alignment rate (or mapping rate):** this describes how many reads, produced by a specific base-caller, can be aligned back to the source reference genome. This rate provides an important metric, because it quantifies how many of the estimated reads possess good enough quality to permit high-level of genome analysis (such as SNP detection). Of course, similar to the base-calling error rate, the alignment rate depends to a large extent on the specific sequence alignment tool that is used. In Table 2, we present the alignment rates for the sequence reads produced by the different base-callers for each of the three datasets. The reads were aligned using Bowtie with conservative parameters (low sensitivity).

It is shown that for all three datasets, a bigger percentage of the sequence reads can be aligned back to the reference if TotalReCaller’s strategy is used for base-calling. Similarly to the base-calling error rate, a higher alignment weight $w_{\text{align}}$ for TotalReCaller directly relates to a higher alignment rate.

**Base-calling speed:** this describes the time needed to perform base-calling for a given dataset. For most base-callers, the total time to perform base-calling can be divided into a training phase and the actual base-calling. During the training phase, a given base-caller computes the parameters for its underlying error models. Similar to Bowtie and Rolexa, TotalReCaller depends on a set of correct reads to compute its error model parameters. These reads are generated by aligning Bustard reads with Bowtie on a reference genome. In addition, TotalReCaller requires the construction of the Burrows-Wheeler transform of the reference genome, which is then used by the base-by-base aligner during base-calling. Table 2 gives a comparison of the base-calling speed of the different base-callers.

It is shown that in comparison to Bowtie, Rolexa and BayesCall, TotalReCaller uses a shorter training phase. The relatively long base-calling time for TotalReCaller can be accounted by the time TotalReCaller implicitly spends on genome alignment while base-calling.

**SNP specificity and sensitivity:** aside from TotalReCaller’s relative performance advantage in terms of error and alignment rates, it may be questioned whether TotalReCaller’s bias due to reference-based Bayesian prior is the source of this advantage, and could affect (perhaps adversely) its single nucleotide polymorphism (SNP) sensitivity and specificity. Specifically, since TotalReCaller injects knowledge from a reference genome into the base-calling process, it is possible that sequence reads at true SNP positions (containing information from positions where the reference genome differs from the genotype that is sequenced) are called incorrectly. Thus, it is vital to examine what happens when a sequence read is called if that sequence contains one or more SNPs with respect to the reference genome. In order to assess this bias toward the reference genome, particularly with respect to reference independent base-callers, we collected the statistics described below and presented the results graphically (Fig. 3). We defined two such important statistics: **specificity**, $\text{SPC}_k$ (also known as true negative rate, TNR) and **sensitivity**, $\text{SNS}_k$ (also known as true positive rate, TPR) for each cycle (BP position) $k$.

$$\text{SPC}_k = \frac{\text{True negatives}_k}{\text{True negatives}_k + \text{False positives}_k} \quad \text{and} \quad \text{SNS}_k = \frac{\text{True positives}_k}{\text{True positives}_k + \text{False negatives}_k}$$

These statistics are based on artificially SNP-inserted reference genomes. On average, we inserted one SNP every $n$ bases randomly.
Table 2. Speed and alignment comparison

<table>
<thead>
<tr>
<th>Genome</th>
<th>Base-caller</th>
<th>ttraining (h)</th>
<th>tcalling</th>
<th>Alignment rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Bustard</td>
<td>~8</td>
<td>~2</td>
<td>15.29</td>
</tr>
<tr>
<td></td>
<td>Ibis</td>
<td>~20</td>
<td>~10</td>
<td>32.66</td>
</tr>
<tr>
<td>poplar</td>
<td>TotalReCaller (w=0.1)</td>
<td>40.50</td>
<td>46.85</td>
<td></td>
</tr>
<tr>
<td>E.coli</td>
<td>TotalReCaller (w=0.3)</td>
<td>~1</td>
<td>77.19</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TotalReCaller (w=0.5)</td>
<td>~1</td>
<td>87.47</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TotalReCaller (w=0.9)</td>
<td>~1</td>
<td>92.89</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bustard</td>
<td>~16</td>
<td>~9</td>
<td>25.55</td>
</tr>
<tr>
<td></td>
<td>Ibis</td>
<td>~16</td>
<td>~9</td>
<td>25.55</td>
</tr>
<tr>
<td>poplar</td>
<td>TotalReCaller (w=0.1)</td>
<td>25.60</td>
<td>25.60</td>
<td></td>
</tr>
<tr>
<td>E.coli</td>
<td>TotalReCaller (w=0.3)</td>
<td>27.74</td>
<td>27.74</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TotalReCaller (w=0.5)</td>
<td>~1.5</td>
<td>29.85</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TotalReCaller (w=0.9)</td>
<td>~1</td>
<td>31.84</td>
<td></td>
</tr>
</tbody>
</table>

In the third column, \( t_{\text{training}} \) tabulates the approximate duration of the training phase, the parameter estimation, for each of the base-callers. In the fourth column, \( t_{\text{calling}} \) tabulates the duration of the actual base-calling. In the last column, ‘alignment rate’ shows the percentage of how many of the reads, called by a specific base-caller, could be aligned back to the reference genome using Bowtie (Langmead et al., 2009).

F. Menges et al.

2334

In the Table 2, the speed of alignment comparison for different base-callers is presented with respect to the E.coli dataset. The TotalReCaller's sensitivity for a low alignment weight is surpassed by Ibis and Bustard for the E.coli dataset. Increasing TotalReCaller's alignment weight increases the specificity and reduces the sensitivity. Considering the significantly higher alignment rate of TotalReCaller (Table 2), the loss of sensitivity from increasing \( w_{\text{align}} \) is at least partially compensated for by the significantly higher alignment rate of TotalReCaller.

The performance of TotalReCaller leads to the conclusion that, by using a Bayesian approach that relies on dynamically creating a reference-based prior, it is possible to significantly lower the error rates of Illumina short reads for both small and large genomes, independent of the sequencing technology used (GAI or GABs). This performance improvement leads directly to higher alignment rates, which in turn implies that more reads from a sequencing run can be used for high-level analysis. The SNP sensitivity and specificity evaluation show that lower error rates and higher alignment rates come at a cost: SNPs can get lost within a single read. However, relating SNP sensitivity and alignment rates it is clear that TotalReCaller represents a significant improvement in the field of base-calling.

METHODS

TotalReCaller combines the knowledge from sequences’ raw intensity data with information from a reference genome. In other words, it generates the most plausible n-base string (out of \( 4^n \) possibilities) that is most likely to have generated the channel intensity data, and also most likely to have originated at some location on the reference genome. The main innovation of TotalReCaller is to tame the worst-case exponential complexity of the implementation by using a branch-and-bound strategy. Specifically, this strategy is used to concurrently extend multiple high-quality reads that are immediately validated not only by the intensity signals, but also by the likely alignments to a reference genome (thus the genome providing a weak prior to a Bayesian inference). This scheme builds on a rigorously defined Bayesian score function that accounts for both, thus resulting in a single score quantifying the quality of a given sequence read. In order to execute this task, TotalReCaller implements four different components that are described in detail in the following sections: (i) linear error model; (ii) base-by-base sequence alignment; (iii) branch-and-bound read extension; and finally (iv) score function.

2.1 Linear error model and filter

We devise a simple linear model to correct errors resulting from cross-talk, fading and cycle-dependent synchronous-lagging. The model is based on a cycle-dependent transition matrix (thus dynamic) in order to filter the raw intensity channels.

We first derive a linear-algebraic model for cross-talk, fading and cycle-dependent synchronous-lagging. The model is based on a cycle-dependent transition matrix (thus dynamic) in order to filter the raw intensity channels.

Let \( I_k \) be the vector of the four raw intensity channels. In order to model cross-talk in cycle \( k \in \mathbb{N} \), we introduce the cross-talk matrix \( A_k \in \mathbb{R}^{4 \times 4} \) and the cross-talk-free channels \( X_k = (X_{k,1} \ X_{k,2} \ X_{k,3} \ X_{k,4})^{\top} \in \mathbb{R}^4 \). We model their relationship with the following formula:

\[
I_k = A_k \cdot X_k. \quad (1)
\]

Since a separate cross-talk matrix is computed for every cycle \( k \), we implicitly normalize the intensities, thus additionally accounting for fading.

Lagging is then modeled by introducing a dependency between the current cycle and the previous cycle, resulting in:

\[
I_{k-1} = \begin{pmatrix} A_{k-1} & 0 \\ Y_k & A_k \\ X_{k-1} & X_k \end{pmatrix}. \quad (2)
\]
SNP specificity (SPC) measures the rate at which a difference between a read and its reference represents a SNP and not a base-calling error. SNP sensitivity (SNS) measures the rate of called SNPs with respect to all SNPs that should be called. (a) phiX-SPC; (b) E.coli-SPC; (c) poplar-SPC; (d) phiX-SNS; (e) E.coli-SNS; (f) poplar-SNS.

\[
\begin{bmatrix}
X_{k+1} \\
X_k
\end{bmatrix} = \begin{bmatrix} A_{k-1} & 0 \\ Y_k & I_k \end{bmatrix} \begin{bmatrix} I_{k-1} \\
I_k
\end{bmatrix},
\]

(3)

where \( Y_k \in \mathbb{R}^{4 \times 4} \) describes the coupling between \( I_k \) and \( I_{k-1} \). A matrix inversion results in a simple transition matrix \( G_k \), which is then used to filter the raw intensity channels. The elements of the matrices \( Y_k \) and \( A_k \) are obtained by statistical analysis of the intensities, using a library of correct reads similar to the training sets used for the parameter estimation of the support vector machines in Alta-Cyclic and Ibis. However, notice that for TotalReCaller this is not a computationally expensive task since it only solves a simple linear system. After applying the filter to the set of raw intensities (Fig. 1), we were able to significantly improve the quality of the intensity channels, as shown in Figure 4.

The error model and filter can easily be extended to include leading (pre-phasing). Our preliminary results showed that the improvement in quality is only minor compared to the increase in computational cost, so we decided to keep the model simpler and excluded modeling leading.

With the intensity channels suitably filtered, we needed a metric to compare the intensity channels with one another. For this purpose, we focused on the conditional probabilities \( P_k(X_k | B) \) and \( P_k(X_k | \neg B) \) with \( B \in \{A, C, G, T\} \). \( P_k(X_k | B) \) denotes the conditional probability of the filtered intensity \( X_k \) of channel \( B \), given that the correct base to call is base \( B \), whereas \( P_k(X_k | \neg B) \) denotes the conditional probability of the filtered intensity \( X_k \), given that the correct base to call is not base \( B \). Since the filtered channels \( X_k \) are independent of each other, we can approximate these probabilities assuming that they are normally distributed (with subscript \( k \) suppressed to avoid clutter).

\[
X_k | B \sim \mathcal{N}(\mu_B, \sigma_B) \quad \text{and} \quad X_k | \neg B \sim \mathcal{N}(\mu_{\neg B}, \sigma_{\neg B}).
\]

That is:

\[
P_k(X_k | B) = \frac{1}{\sqrt{2\pi\sigma_B}} \exp\left(-\frac{(X_k-\mu_B)^2}{2\sigma_B^2}\right),
\]

(4)

\[
P_k(X_k | \neg B) = \frac{1}{\sqrt{2\pi\sigma_{\neg B}}} \exp\left(-\frac{(X_k-\mu_{\neg B})^2}{2\sigma_{\neg B}^2}\right).
\]

(5)

The means together with their SDs have already been presented in Figure 4, in order to show the workings of the linear error model and filter.

2.2 Base-by-base sequence alignment

The key idea of TotalReCaller is to perform alignment while the sequence is being base-called. The partially generated sequences, which are grown one base at a time, must be aligned back to the reference genome. To account for
Table 3. Example: probabilities from FM search for each base preceded by 'ACGAC'

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Frequency ( f_B )</th>
<th>( P_{\bar{B}} )</th>
<th>( P(B) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACGACA</td>
<td>100</td>
<td>0.10</td>
<td>0.90</td>
</tr>
<tr>
<td>ACGACC</td>
<td>20</td>
<td>0.02</td>
<td>0.98</td>
</tr>
<tr>
<td>ACGAG</td>
<td>500</td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>ACGACT</td>
<td>380</td>
<td>0.38</td>
<td>0.62</td>
</tr>
</tbody>
</table>

this computationally intensive task, we designed an efficient base-by-base aligner that is based on a suffix tree search algorithm. Inspired by the many Burrows-Wheeler based short read sequence aligners (Bowtie (Langmead et al., 2009); SOAP2 (Li et al., 2009); BWA (Li and Durbin, 2009)), we constructed our base-by-base essentially on the same principles, specifically the Ferragina-Manzini (FM) search algorithm (Ferragina and Manzini, 2000) and the Burrows–Wheeler transformation (BWT) (Burrows and Wheeler, 1994).

Ferragina and Manzini showed how the suffix tree of a reference genome can be accessed through its BWT, which does not require more memory than the reference genome itself. Thus, searching for a (partial) sequence read in a BWT reference can be performed highly efficiently. In addition, not only information about the existence but also the number of occurrences in the given reference of the (partial) sequence read can also be computed concurrently. For more details about the FM search algorithm, see the related publication (Ferragina and Manzini, 2000).

Note that, in general, sequence aligners are usually only interested in whether and where a sequence read is located in a given reference. In contrast, building solely on the alignment information for base-calling, it suffices for us to retain only the frequency of a partial sequence read. Consider the example that the (partial) sequence ‘ACGAC’ is contained in a reference genome. We can use the FM search to count how often the sequence ‘ACGACB’ with \( B \in \{A, C, G, T\} \) are contained in the reference, from which we can then compute the probability \( P_B(x) \), that the next base (at cycle \( k \)) in the sequence is \( B \):

\[
P_B(x) = \frac{f_B}{f + f_A + f_C + f_T} \quad \& \quad P(x \neq B) = 1 - P(B)
\]

Table 3 shows a complete example of how the base probabilities are computed using the FM search. Now that we have introduced the intensity filter and base-by-base alignment components, what remains to be shown is the branch-and-bound scheme (Land and Doig, 1960; Lawler and Wood, 1966), the worst-case complexity is rarely encountered in practice (and can be further controlled by beam-search). Note that the special situations in which the exponential worst-case behavior would be exhibited occur when the sequence is extremely noisy and/or when the reference is incorrect (or highly error-ridden), thus producing exponentially more plausible hypothetical sequence reads; a judicious solution in these cases would then involve terminating the sequence read at a smaller read length or rejecting it outright.

The algorithm can now be described as a sequence of three consecutive steps that are repeated once for each cycle:

1. **Branching:** for each sequence in the solution space \( N_{k-1} \), all four possible successor sequences are generated, resulting in the solution space \( N_k \). Note that at this point \( N_{k-1} \subseteq N_k \).

2. **Bounding:** each sequence in \( N_k \) is evaluated according to a score function \( f_{score} \). The score \( s_k \) of a specific sequence at cycle \( k \) is updated as follows:

\[
s_k = s_{k-1} + f_{score}
\]

3. **Pruning:** all but the best (highest score) \( k \in N \) sequences are pruned, thus reducing the size of \( N_k \) to \( |N_k| = 1 \).

Note that, by reducing the computational complexity through bounding the solution space, we are not guaranteed anymore to generate the optimal solution. However, in practice the accuracy of the algorithm’s outputs is only slightly affected. Where necessary, the computational cost can be traded off for higher accuracy by setting a parameter that controls the beam-width of a beam-search.

### 2.4 Score function

To complete the description of our base-caller, we need to define the score function \( f_{score} \), used to evaluate the quality of the candidate sequences in the tree. Using Bayes’ theorem, we estimate the probability \( P_B \) that a specific base \( B \in \{A, C, G, T\} \) is indeed the correct base to call at cycle \( k \), given the filtered intensity vector \( X_k \), as:

\[
P_B(X_k) = \frac{P(X_k | B) \cdot P(B)}{P(X_k)}
\]

Since for our purposes it is sufficient to have a quantitative measurement (not a probability) to compare all different solution to one another, we introduce a simplified score function \( f_{score} \) which is based on \( P_B(X_k) \):

\[
f_{score} = \log \left( \frac{P_B(X_k | B)}{P_B(X_k | \bar{B})} \right) = \log \left( \frac{P_B(B)}{P_B(\bar{B})} \right) + \log \left( \frac{P_B(X_k | B)}{P_B(X_k | \bar{B})} \right)
\]

Therefore, the score function consists of two parts, both of which can be computed independently according to the sections discussing the intensity filter and the base-by-base alignment algorithm. The weight \( \alpha \) allows for a user-defined control over the impact of the alignment on the overall score, thus enabling the user to adjust the Bayesian bias, appropriate for a particular application.

### 3 DISCUSSION

This article introduces a new base-caller, TOTALReCaller, which opens new avenues for reducing errors significantly in...
short sequencing reads simply by injecting knowledge from a reference source genome through a base-by-base alignment algorithm. Stepping back and re-examining the overall strategies for base-calling, alignment, resequencing, polymorphism-detection, sequence assembly and assembly validation, it appears that there is much to be gained from integrating various steps in genomics analysis that have been traditionally performed sequentially. A Bayesian framework can be developed rigorously for the purpose of such integration, and classical branch-and-bound algorithms can be used to tame the efficiency, which otherwise could make the integration intractable.

In addition to a software implementation of TotalReCaller, we have also implemented and validated a proof-of-concept Field-Programmable-Gate-Array (FPGA) design. Our hardware design is capable of performing all parts of the algorithm, presented here, including: filtering the raw intensity channels, FM search, score function calculation and the branch-and-bound algorithm. Building upon this FPGA design, we wish to next show that re-sequencing function calculation and the branch-and-bound algorithm. Building upon this FPGA design, we wish to next show that re-sequencing can be performed in an embedded environment. Coupled with recent advances in sequencing chemistry (e.g. rapid field-effect-transistor-based assay of pH changes after a base incorporation by a polymerase), this process could pave the way for a real-time SNP-calling machine.

To move further in this direction and to fully exhaust TotalReCaller’s potential, much remains to be done. For instance, we will next need to compare TotalReCaller to other sequence aligners. This article primarily focused on TotalReCaller’s base-calling qualities, describing the algorithm and comparing it to its peers. As our analysis begins to focus on sequence alignment (e.g. including gapped alignment), we expect to better understand how TotalReCaller handles various structurally complex portions of the genome that often frustrate most suffix tree-based aligners.

As discussed earlier, another direction of research currently focuses on developing a non-parametric approach to handle intensity errors—thus, without having to learn optimal strategies from a calibrating genome (or training data). Currently, TotalReCaller (like all other parametric base-calls) depends on a training set of correct reads and runs the risk of being misled, should the training data be corrupted by some fraction of incorrect reads. Last but not least it will be important to study the effects bounding onto the the overall performance. The results presented in this article were generated with a static bound of $l = 32$. A static bound was chosen in order to allow an efficient low-level hardware implementation. The value $l = 32$ was empirically chosen. Future improvements of TotalReCaller’s software implementation will include a dynamic bound, which has the potential to improve both accuracy and/or speed.

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**REFERENCES**


