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e-RGA: enhanced Reference Guided Assembly of Complex Genomes



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Abstract

Next Generation Sequencing has totally changed genomics: we are able to produce huge amounts of data at an incredibly low cost compared to Sanger sequencing. Despite this, some old problems have become even more difficult, *de novo* assembly being on top of this list. Despite efforts to design tools able to assemble, *de novo*, an organism sequenced with short reads, the results are still far from those achievable with long reads. In this paper, we propose a novel method that aims to improve *de novo* assembly in the presence of a closely related reference. The idea is to combine *de novo* and reference-guided assembly in order to obtain enhanced results.

Introduction

DNA sequencing is becoming cheaper every day. Instruments like the Illumina HiSeq 2000 or Solid 4 System are able to produce higher than 30X coverage of a human genome for less than \$10,000. Next Generation Sequencing (NGS) allows sequencing at low cost and at a fraction of the time with respect to the Sanger Method [1]. NGS technologies (Illumina, Roche and Solid, to mention just a few) are capable of producing an enormous amount of raw data (for a complete review, see [2]). The throughput of such instruments is increasing so fast that descriptions of their performance became obsolete in just a few months.

Even though many papers have presented high-quality assemblies based on NGS data (see [3,4]), *de novo* assembly, especially for large-genome eukaryota, is still a 'holy grail' [5].

When a completely new organism is to be sequenced, the basic assembly strategy is still the celebrated Whole-Genome Shotgun (WGS) method. A number of tools that aim to perform de novo assembly using NGS data have been proposed. Among the most popular, we mention SOAPdenovo [6] and ABySS [7] (see [8] for an updated list). The assemblies produced by tools designed for NGS data are, in general, not comparable in quality with the assemblies produced by instruments like Arachne [9] and those designed for capillary sequencing data. The reason for this is that while, with NGS instruments, coverage is no more a bottleneck, read length, as well as the reduced size of the insertion between paired reads, makes correct assembly and positioning of repeats much more of an issue. De novo assembly with short reads is still very difficult [5] and, when assembling complex and repeated genomes, reasonably conservative de novo assembly programs are likely to produce collections of highly fragmented contigs. An interesting strategy to improve de novo assemblies has been termed "assembly reconciliation". The goal of assembly reconciliation is to merge the assemblies produced by different tools and to detect possible mis-assemblies [10].

The number of organisms whose genomes have been completely sequenced has been increasing rapidly each year and, for this reason, it is becoming viable to sequence an organism and then to align the sequence against a closely related genome. This strategy goes by the name of Reference Guided Assembly (RGA), its main advantage being the fact that, in general, even low coverage is sufficient to yield useful results. RGA consists of two phases: first, all the reads are aligned against the reference genome; then, a consensus sequence is extrapolated. Everywhere the coverage drops to zero, a sequence of Ns is placed. This problem has already been studied in the context of Sanger sequencing. In [11] and [12], two methods were proposed that use reference sequences to assist the assembly of new organisms. The challenge is becoming even more interesting with the advent of next generation sequencers, beginning with the technicalities and the practical considerations involved in the alignment phase.

As a matter of fact, many tools capable of rapidly aligning millions of short reads against a reference genome have been proposed recently. Tools like SOAP2 [13] and rNA [14] are essential for performing reference-guided assembly. The main problem with RGA and NGS is that (essentially for efficiency reasons) mapping al-

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gorithms are highly conservative: it is possible to align reads with only a low number of errors and usually without gaps. In other words, we are able to reconstruct the conserved regions, while we cannot reconstruct areas that are divergent and (usually) more interesting. While, for example, there are techniques that use the paired-reads information to identify insertions/deletions (structural variations) [15,16], there is no clear way to reconstruct them.

In [17], a tool (MAIA) has been proposed to integrate multiple *de novo* and reference-guided assemblies. This tool uses the output of different assembles, and of different reference-guided assemblies obtained with several reference sequences, to improve the final assembly result. MAIA constructs an overlap graph from the pairwise alignments of all the contigs. In large and repetitive genomes, like plant genomes, this step is computationally expensive and could easily lead to a large number of ambiguous or false overlaps.

Velvet's Columbus module tries to improve the assembly results using a reference sequence. In particular, the Columbus module aims to reconstruct candidate structural variations. Again, in this case, as a consequence of the repeats, this approach cannot be used on large, repetitive genomes.

In this paper, we briefly discuss and present results of a novel strategy to assemble genomes in the presence of a related sequence. In particular, we study how to merge the de novo and reference-guided assembly strategies, in order to assemble new organisms and improve the result achievable using only either one of the two. We will show how, by applying some of the ideas of assembly reconciliation, we can obtain an enhanced reference assembly of a new organism. All the alignments are guided by the reference sequence, in this way avoiding mis-assemblies and ambiguous overlaps. In [18], we illustrated the results obtained by our pipeline on a set of small organisms (chloroplasts and microbial); in this work, we show how the same pipeline can be effectively used for the assembly of large, highly repetitive and heterozygous genomes (plant genomes).

Reference-guided assembly

When a reference sequence A and a set of reads R are given, there are essentially two possible

ways to perform reference assembly. The standard way simply involves aligning all the reads in R against the reference A, and then obtaining some consensus sequences. In the text that follows, we refer to this method as standard-RGA (s-RGA), and similarly, we call the consensus sequence produced s-A. An alternative approach is to perform de novo assembly on R first, and then to align the resulting contigs against the reference A. We call this second method de novo-RGA (dn-RGA), and the consensus sequence produced from it dn-A. Both the output sequences have "N" everywhere the coverage drops to 0. In order to simplify the discussion, we suppose A to be a single sequence (and therefore s-A and dn-A are also single sequences). It will be clear that this is not a limitation.

In the presence of NGS data, we have to use aligners like SOAP2 [13] and rNA [14] to obtain s-A. These aligners are highly conservative, they allow alignment of reads with a low number of errors, usually without gaps. For this reason, the length of s-A is the same of A. The sequence dn-A is obtained in three phases: first, reads are assembled using a short-read assembler ([6,7]); the resulting contigs are then aligned against A; after this, the consensus is generated. The challenge is to find an order for the contigs generated through the de novo assembly procedure. Several tools have been proposed to address this task. OSLay [19] computes a synthetic layout of the contias using a reference sequence to anchor the de novo sequences; the Mauve aligner [20] gives as output an ordered version of the *de novo* contigs; and PGA [21] is able to layout the contigs with more than one reference genome at a time using global searches. All these tools implement or use a BLAST-like [22] search to align contigs against the reference. This alignment technique allows us to place reads on a reference with low similarity constraints. In particular, the contias can be aligned against the reference sequence allowing partial hits and gaps.

This situation is similar to the already studied situation of assembly reconciliation. Casagrande et al. [10] proposed a method capable of merging two draft assemblies without performing global alignment. In particular, they proposed the use of one of the two assemblies as an "anchor", in order to resolve conflicts (Master Assembly).

Given the two assemblies s-A and *dn*-A, a suitable adaptation of this idea can be applied

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in the current context to obtain an enhanced reference assembly.

Methods

The Merge Graph: Definition

In this section, we briefly sketch the formal steps necessary to define the Merge Graph, at the base of our technique. More details can be found in [18].

With S[i,j] (S being either Δ or Γ), we identify the so-called "slice" of a string S, namely the substring of S from position i to j. If S[i,j] belongs to $\{a,c,g,t\}^*$, we call it a (pure) contig, while if S[i,j]belongs to $\{N\}^*$, it is named gap. In this context, when a (pure) contig is maximally extended, we say that it is a max-contig, and we define, in an analogous way, a max-gap (in general, we speak of max-area). Given two strings δ and γ , the function $D(\delta_{\gamma,d})$ returns a value between 0 and 1, representing a percentage difference between the two strings. This value is naturally computed using a distance metric d (e.g., Hamming). The merge graph $MG(\Delta,\Gamma)$ is a directed graph such that V is contained in $I_{\Delta} \times I_{\Gamma}$ (I_{Δ} and \tilde{I}_{Γ} being all possible intervals in \varDelta and \varGamma , respectively) and can be partitioned into four sets: gap-nodes (V_{α} , gap nect pairs of intervals (one on Δ and one on Γ) against gap), delta-nodes (V_s, a Δ -contig against that can subsequently be put in the output as-

idble 1.	Table of symbols and definitions.				
А	Reference sequence				
s-A	Consensus sequence obtained aligning short reads against refer- ence A				
dn-A	Consensus sequences obtained aligning de novo contigs against ref- erence A				
e-A	Enhanced reference-guided as- sembly obtained through the e-RGA pipeline				
Δ, Γ	Generic sequences				
$MG(\Delta,\Gamma)$	Merge graph for the sequences \varDelta and \varGamma				
MGA	Merging Global Alignment				

a Γ -gap), gamma-nodes (V,, a Δ -gap against a Γ -contig) and merge-nodes (V_m, contig against contig). We also fix t and s, two thresholds that express bounds on the distance (alignment similarity) and the absolute position (within the relative string), respectively.

Edges are defined in such a way as to con-

11 21 31 41 51 61 Δ : NNNNNNaag gtttaaggtc ctacaNNNNa ctcatcataa aaacccNNNN NNNNNctct aggtaaaa Γ : NNNNNNNNN NNNNggccta cacNNNNNac tcatcatcaa aaacccNNaa aaaaNNNNNN NNNNaaaaa

Figure 1 (a). The strings Δ and Γ ($|\Delta|$ =68, $|\Gamma|$ =69).



Figure 1 (b). A possible Merge Graph G for Δ and Γ with s=2 and t=2.

 Δ' : NNNNNNN AAGGTTTAA GGTCCTACA- NNNN- ACTCATCATAAAAA-CCC NN NNNNNN NN- CTCTAGG TAAAA Γ' : NNNNN-- NNNNNNNN GG-CCTACAC NNNNN ACTCATCAAAAAACCC NN AAAAAA NNN NNNNNN AAAAA

 Λ :

Figure 1 (c). A possible Global Alignment obtained from the Merge Graph G (M means match, S means substitution, I means insertion, while D means deletion).

NNNNNNN AAGGTT- TAAGGTCCTACA- NNNN- ACTCATCAT-AAAAAACCC NN NNNNNN NN- CTCTAGG TAAAA Λ' : Γ' : NNNNNNN NNNNNNN ---GG-CCTACAC NNNNN ACTCATCAACAAAACCC NN AAAAAA NNN NNNNNNN AAAAA constrain violated

Figure 1 (d). A Global Alignment that does not allow the creation of an MGA.



Figure 2. MGA construction. A possible scenario.

In Figure 1B, an example of a merge graph is given.

In order to clarify the notation, we have summarised the symbols used throughout the paper in Table 1.

Merge Graph and Global Alignment

Given the strings Δ and Γ , there is a deep connection between the merge graph $MG(\Delta,\Gamma)$ and a global alignment between them. Building global alignments turns out to be equivalent to building a merge graph. In following text, we refer to the global alignment ensuing from a merge graph as Merging Global Alignment (MGA).

The merge graph $MG(\Delta,\Gamma)$ can be used to extract a family of edit strings. From each node, we can produce an edited version of the two substrings that are represented. From delta and gamma nodes, we simply extract the contigs, while from the merge and gap nodes we can produce an edited version of one of the two strings. In the case of gap nodes, the edited version will contain only insertions and deletions (see Figures 1A, 1B and 1C). Once the edit strings are computed, the corresponding MGA can be calculated.

Given an MGA, the construction of a merge graph is more complicated. An MGA is a global alignment with two kinds of local properties: locality and similarity. In general, a global alignment does not guarantee these local properties, hence we can easily construct a global alignment that violates a local constraint. It can be proved that the global alignments we are seeking must respect the following properties: if $\Delta[i,j]$ and $\Gamma[k,l]$ are aligned one against the other, then |i-k|-1 < s and |j-1|-1 < s (locality); if $\Delta[i,j]$ and $\Gamma[k, l]$ are aligned and at least one is a max-contig or $\Delta[i-1] = \Gamma[l+1] = N$ (or $\Delta[j+1] = \Gamma[k-1] = N$), then $D(\Delta[i,j],\Gamma[k,l],d)$ (similarity).

If such an alignment exists, calling \varDelta' and \varGamma' the two strings over the alphabet $\{a, c, g, t, N, -\}$ returned by the global alignment between \varDelta and Γ , we can build $MG(\Delta,\Gamma)$ by simply reading from left to right Δ' and Γ' . For each position, we have

sembly. The resulting graph must be acyclic. to judge if a new node is beginning, or if we can continue extending the current one.

> The determination of this global alignment can be computationally cumbersome. We cannot simply use an algorithm that calculates an optimal sequence alignment because a choice that can create an optimal global alignment will not necessarily lead to an alignment that respects all the local constraints (see Figure 1(d)). We will now sketch a complete algorithm that, given the strings \varDelta and Γ , generates all possible $MG(\Delta,\Gamma).$

> The algorithm starts by reading the two sequences from left to right. For every contig in \varDelta and Γ_{i} we can recursively compute all the possible alignments that satisfy the locality, and possibly the similarity, constraints. More detail is given in Figure 2. Let us assume that the last generated node was $\langle z, i \rangle, [m, v] \rangle$. From the merge graph definition, we have that at least one of i or v must be the end of a max-area, in this case i. At this point, we have to calculate the nearest (from i) max area end, *n* in the case shown in Figure 2. So the node we are going to create is $\langle i,k \rangle, \langle v,n \rangle \rangle$, with n - a < k < n + s (paying attention to some special case, we can reduce the search space). In order to generate all the possible graphs, we have to recursively generate all the nodes. In case we are generating a merge node, we have also to check if the similarity constraint is respected. The algorithm terminates because at every step it proceeds forward along both strings.

Minimal Merge Graph

Given two strings Δ and Γ , it is clear that the existence of an $MG(\Delta,\Gamma)$ depends on the two thresholds s (the bound on the relative distance between intervals involved in the same node) and t (the bound on the similarity distance between strings involved in the same merge node). By setting s to be large enough, we can easily go towards computing a merge graph composed only by gap, delta and gamma nodes. Another trivial solution can be found when t is set in such a way that merge nodes accept very low levels of similarity.



Figure 3. Practical identification of Merge-nodes for the Merge Graph Construction.

It is clear that a merge graph between a pair of sequences is interesting when s is a small constant when compared to Δ and Γ lengths and when t is sufficiently strict. These two constraints can strongly help in designing a better performing algorithm.

Merge graph and referenceguided assembly

A merge graph is a data-structure able to describe a global alignment between two strings, with further constraints on local alignment and similarity. This data-structure can be used both to describe the relations between two strings, and to extract a consensus.

When working with s-A and dn-A, we elect one of the two sequences to be the Master Assembly (MA), that is the assembly we believe to be correct. In practice, in presence of a merge node, instead of calculating a consensus, we simply keep the sequence from the MA. Usually, even though two choices are possible, the MA will almost always be dn-A, with regions present in the sequenced organism and absent in the reference. If the merge graph MG(s-A,dn-A) is available, it can be used to extract a new assembly. Each node p in MG(s-A, dn-A) is characterised by two intervals, [i,j] and [k,l]. For $p = \langle [i,j], [k,l] \rangle$, we extract the sequence dn-A[k, I], if p is a gamma or a merge node, s-A[i,j], if p is a delta-node, or the shortest between s-A[i,j] and dn-A[k,l], if p is a gap node. This assembly is named e-A (enhanced Reference Guided Assembly).

The difficult part is the MG(s-A, dn-A) construction. The correct and complete algorithm presented in section 3.2 takes a time proportional to $O(s(j-i)^2)$ for each max-contig in the worst case. Additionally, the parameters *s* and *t* are unknown and, in general, there is no clear way to estimate them in advance, or to at least sensibly approximate them.

When working with s-A and dn-A, we have that the MG(s-A,dn-A) merge graph must exist for some s belonging to the set $\{0,...,|dn$ -A|-|s-A|}. This follows directly from the construction of the two strings. It is more difficult to limit t. A good working approximation is the percentage difference allowed in the de novo contig alignment.

The particular context provided by *s*-*A* and *dn*-*A* allows us to further improve the construction algorithm, concentrating only on a significant subset of all the global alignments associated to *MG(s*-*A*,*dn*-*A*). Thanks to some intrinsic properties of *s*-*A* and *dn*-*A*, the brute-force algorithm can be improved, avoiding the generation of all possible global alignments. See Figure 3 for a graphical representation, and [18] for further details.

Enhanced-rga: implementation details

The pipeline represented in Figure 4 was implemented using several third-party tools and a set of Perl scripts implemented by the authors.

In order to construct s-A, first, a short-string aligner is used to align all the reads against the reference sequence, and a consensus is then extracted. In all cases in which a read is found in multiple occurrences, we randomly choose one of the alignments. We used the short-string aligner rNA [14] and the "pileup" command provided by samtools [23] to extract the consensus sequence.

dn-A was obtained by first performing *de novo* assembly with ABySS [7] and the CLC assembler Cell 3.0 [24]. Together with SOAPdenovo, [6] these are the only two assemblers able to assemble complex genomes using a reasonable amount of time and RAM memory. We noticed that, using contigs from both assemblies, the amount of genome reconstructed in *dn-A* greatly improves. The delicate phase of mapping contigs against

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Figure 4. eRGA implementation

the reference sequence was accomplished with the <u>CLC-Workbench</u>¹.

Although we used these specific tools, clearly the production of *s*-*A* and *dn*-*A* can be carried out using different software without significant modification of the pipeline.

The core of e-RGA is the MG(s-A,dn-A) construction and e-A generation. These crucial phases are implemented within a Perl script that uses BLAST [22] to perform the approximate alignment. The program first memorises both s-A and *dn-A*, and localises all the max-areas (max-contigs and max-gaps). The MG(s-A,dn-A) construction proceeds as outlined in Section 4, with all the alignments performed by BLAST.

The software, together with a small example, can be downloaded at <u>http://sole.dimi.uniud.</u> <u>it/~francesco.vezzi/software.php</u>.

Experiments and results

The Datasets

In [18], the e-RGA pipeline was tested on small genomes, demonstrating the effectiveness of our pipeline. We have further improved our pipeline and successfully used e-RGA on two large and complex plant genomes. The first dataset, named Sangiovese, comprises 5 Illumina lanes used to re-sequence a grapevine variety (Sangiovese, Rauscedo clone R24) with 100bp paired-end reads for a 90X total raw coverage. For this dataset, we used as reference sequence the genome of the highly homozygous grape clone, PN40024, used as reference genotype from the French-Italian Consortium for grape genome characterisation [25]. The second dataset, named Poplar, comprises 6 Illumina lanes used to sequence a Poplar individual belonging to the Populus nigra species, with 100bp paired-end reads for an 85X total raw coverage. In this case, we used as reference sequence the Populus trichocarpa genome [26]. While in the Sangiovese dataset we used a reference sequence belonaing to the grapevine species, in the Poplar dataset the reference belongs to a different species. This difference is important in order to understand the differences obtained in the results. In both cases, before assembling and aligning, all the reads were filtered for quality, and we eliminated all sequences belonging to chloroplasts and mitochondria.

Both grapevine and poplar are characterised by long, repetitive genomes (480Mbp and 417Mbp respectively); moreover, both the sequenced individuals are highly heterozygous. These three conditions (length, repetitiveness and heterozygosity), together with the presence of two reference genomes, are perfect for our pipeline.

Results Discussion

Tables 2 and 3 summarise the results from the Sangiovese and Poplar datasets. As a measure of the assembly quality and correctness, we report the percentage of aligned reads (the same reads used to perform reference and *de novo* assembly), the number of contigs reconstructed, the mean contig length, the L50g (the length of the longest contig such that the sum of all the contigs greater than it represents half the expected genome length) and, in brackets, the L50c (the length of the longest contig such that the

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sum of all the contigs greater than it represents half the total contig length), and the percentage of Ns in the sequence. The L50g gives us a normalised value that describes the connectivity level of the assembly.

Those statistics have been computed for the reference sequence A, for the s-RGA output s-A, for the de novo assembly output dn, for the dn-RGA output dn-A, and finally, for the e-RGA output e-A.

Statistics such as the mean contig length, contig number, L50c and L50g, give us an idea of the quality of the assembly. In the Sangiovese case, we can see how the mean length obtained through e-RGA is longer than the other approaches, and although the dn-A mean length has a close value, we must consider the fact that these contigs cover only half the genome length as described by the high percentage of unknown characters. In the Sangiovese dataset, the most impressive results are the L50g and L50c improvements. Both e-A's L50g and L50c are better than those of s-A, and they largely improve the results achievable with de novo assembly alone. This shows that our pipeline can effectively improve the final assembly result.

the Poplar dataset. Owing to the distance be-Table 2. Results obtained for the Sangiovese dataset.

tween the sequenced organism (Populus nigra) and the reference genome (Populus trichocarpa), the Poplar results can look less promising than those of Sangiovese. However, the number of mapped reads against e-A is higher than the number of reads mapped against both s-A and dn-A. The fact that we are able to map a higher number of reads against dn should also be a consequence of the distance between the reference and the sequenced genome. As far as the standard assembly statistics are concerned (L50g, L50c and mean contig length), we can again see how the e-A results are better than those achievable by simply mapping reads or contigs back to the reference. Despite the de novo assembly result looking much better than the other approaches, we must stress the fact that de novo assembly alone gives us a set of 289,854 unordered contigs, with no information about their position in the final genome. It would be interesting, but outside the scope of this work, to understand the composition of the contigs not used to construct dn-A. These contigs, if correctly assembled, represent the areas belonging to the sequenced organism exclusively.

A further measure of the improvements intro-Similar results are summarised in Table 2 for duced by the use of e-RGA is the number of successfully aligned paired reads (i.e., paired reads

For all the techniques used, we show the percentage of aligned reads, the number of contigs, the mean contig length, the L50 length computed both on the expected genome length and on the total contig length, and the number of unknown characters "N".

	Sangiovese						
	% aligned reads	Contigs number	Mean contig length	L50g (L50c)	% Ns		
А	80.21%	-	-	-	3.00%		
s-A	80.99%	246752	1758 bp	8514 bp (9901 bp)	7.64%		
dn	53.10%	289854	1942 bp	1753 bp (3328 bp)	0.70%		
dn-A	50.71%	109833	2246 bp	600 bp (3947 bp)	47.70%		
e-A	81.77%	198194	2282 bp	12494 bp (14219 bp)	6.40%		

Table 2. Results obtained for the Poplar dataset.

For all the techniques used, we show the percentage of aligned reads, the number of contigs, the mean contig length, the L50 length computed both on the expected genome length and on the total contig length, and the number of unknown characters "N".

	Poplar						
	% aligned reads	Contigs number	Mean contig length	L50g (L50c)	% Ns		
А	55.00%	-	-	-	2.14%		
s-A	58.00%	778065	365 bp	525 bp (1105 bp)	25.22%		
dn	67.84%	116683	2728 bp	2906 bp (4487 bp)	0.40%		
dn-A	37.00%	77370	1335 bp	0 bp (2085 bp)	62.46%		
e-A	59.00%	558762	482 bp	957 bp (1959 bp)	18.56%		

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that align on the sequence at the expected distance and orientation). In both datasets, e-A is the sequence on which the largest number of constraints is respected.

More Applications

A possible e-RGA application, not explored in this work, is the identification and, more importantly, the reconstruction of structural variation. Two different steps of the e-RGA pipeline can be instrumental to this purpose. First is in the construction of *dn-A*. We can identify contigs that are aligned with gaps: alignments that introduce gaps in the reference sequence represent a putative insertion in the sequenced genome; conversely, alignments that introduce gaps in contigs reveal a putative deletion in the sequenced genome. The second step in which we might be able to identify structural variations is in e-A construction from the MG(s-A,dn-A) graph. In this case, a gamma-node <[i,j],[k,l]> (dn-A-contig against a s-A-gap) in which the interval [i,j] (the s-A-gap) is shorter than [k,I] (the dn-A-contig) is witness to an 6. Li R, Zhu H, Ruan J al. (2010) De novo asseminsertion in the sequenced genome; conversely, if the interval [i,j] is longer than [k,l] then the node is witness to a deletion in the sequenced genome. In the case of *delta-nodes*, the situation is symmetric.

Conclusions

The e-RGA pipeline was successfully applied to small organisms in [18]; in this paper, we have shown how the same approach can easily scale to large datasets with high coverage over complex (large and highly repetitive) genomes like the Grapevine and Poplar genomes.

e-RGA needs a reference genome belonging to a closely related organism. With the number of available genomes growing at a speed believed impossible only few years ago, this requirement is becoming standard.

Several research efforts are ongoing to design tools to identify and study structural variations (SV) [16], in particular within individual genomes. One major stumbling block is that it is still unclear how the identified or putative SV can be reconstructed. A future development of e-RGA will be to output putative SV identified during e-A construction. In this way, our pipeline, coupled with a tool able to identify/verify SV, could be used to reconstruct sequences that are specific to a particular individual.

Competing interest statement

None declared

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