HINGE: long-read assembly achieves optimal repeat resolution

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Long-read sequencing technologies have the potential to produce gold-standard de novo genome assemblies, but fully exploiting error-prone reads to resolve repeats remains a challenge. Aggressive approaches to repeat resolution often produce misassemblies, and conservative approaches lead to unnecessary fragmentation. We present HINGE, an assembler that seeks to achieve optimal repeat resolution by distinguishing repeats that can be resolved given the data from those that cannot. This is accomplished by adding "hinges" to reads for constructing an overlap graph where only unresolvable repeats are merged. As a result, HINGE combines the error resilience of overlap-based assemblers with repeat-resolution capabilities of de Bruijn graph assemblers. HINGE was evaluated on the long-read bacterial data sets from the NCTC project. HINGE produces more finished assemblies than Miniasm and the manual pipeline of NCTC based on the HGAP assembler and Circlator. HINGE also allows us to identify 40 data sets where unresolvable repeats prevent the reliable construction of a unique finished assembly. In these cases, HINGE outputs a visually interpretable assembly graph that encodes all possible finished assemblies consistent with the reads, while other approaches such as the NCTC pipeline and FALCON either fragment the assembly or resolve the ambiguity arbitrarily.

[Supplemental material is available for this article.]

While genome assembly has been a central task in computational biology for decades, only with the recent advent of long-read technologies has the goal of obtaining near-finished assemblies in an automated fashion become within reach. However, extracting the information present in long error-prone reads in order to reliably resolve repeats is still a challenge (Myers 2016a). Attempts to resolve repeats that are fundamentally unresolvable from the reads at hand—a practice that can be driven by the prospect of a higher N50 score—can lead to incorrect assemblies and ultimately impact downstream scientific analyses. On the other hand, a conservative approach that breaks the assembly at points of seeming ambiguity may fail to produce the longest contigs that can be constructed given the data.

In this sense, an optimal assembler should be one capable of identifying and resolving all, and only those repeat patterns that are resolvable given the available read data. Equivalently, this objective can be viewed as the construction of an *assembly graph* with the maximum level of repeat resolution that is possible given the data. If a finished assembly of the genome is possible, such a graph would consist of a single cycle (in the case of a single circular chromosome). Otherwise, the next-best objective would be the construction of a *repeat graph* (Pevzner and Tang 2001; Mulyukov and Pevzner 2002) where long repeats are collapsed into a single path. Such paths capture inherent ambiguities about the target genome that cannot be resolved given the data. Thus, constructing the *maximally resolved assembly* graph corresponds to minimizing the number of repeat-induced collapsed segments.

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Article published online before print. Article, supplemental material, and publication date are at http://www.genome.org/cgi/doi/10.1101/gr.216465.116. Freely available online through the *Genome Research* Open Access option. As a prerequisite to this task, one must first understand which repeat patterns can be reliably resolved given the set of reads. Early studies of this fundamental problem appeared in the context of sequencing by hybridization (Ukkonen 1992; Pevzner 1995) and were later extended to shotgun sequencing through the notion of bridging (Bresler et al. 2013). A repeat is said to be *bridged* if at least one read completely contains one of its copies (throughout the paper, we use the word copies to refer to the distinct occurrences of a repeat element). The notion of bridging allows us to define a maximally resolved assembly graph as the graph where only segments corresponding to *unbridged* repeats are collapsed, as discussed in Supplemental Figure S1. The de novo construction of such a graph yields the longest contigs that can be reliably constructed, and also describes the plausible arrangements of these contigs in the target genome.

Assembly graphs have been a key component in assembly pipelines since the early days of sequencing projects (Myers et al. 2000). Approaches to assembly graph construction are customarily divided into two categories: de Bruijn graph-based approaches, and overlap-layout-consensus (OLC) approaches. In the de Bruijn framework (Pevzner and Tang 2001; Mulyukov and Pevzner 2002), the set of all *k*-mers is extracted from the reads and used to build a graph where two *k*-mers that appear consecutively in a read are connected by an edge. This construction has the desirable property that the resulting graph is essentially Eulerian, and repeats longer than *k* base pairs are naturally collapsed into a single path. Furthermore, the graph construction is typically followed by repeat resolution steps using reads that bridge repeats. This allows several de Bruijn graph-based assemblers to produce a maximally resolved assembly graph where only unbridged repeats remain collapsed

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Figure 1. The goal of HINGE is to produce a maximally resolved assembly graph, where repeats that are bridged by the reads are not collapsed, and repeats that are unbridged are collapsed in a natural way, similar to what is achieved with de Bruijn graphs. (*A*) If at least one of the two copies of a repeat is bridged (green segments), the maximally resolved assembly graph should separate the two copies. In (*B*–*E*), respectively, we illustrate an unbridged repeat, an unbridged inverted (i.e., reverse-complemented) repeat, an unbridged triple repeat, and a single-bridged triple repeat, and the assembly graph obtained by collapsing segments corresponding to unbridged repeats. Notice that in *B*,*E*, the graph admits a single traversal and can be further resolved, while in *C*,*D*, the graph admits two distinct traversals and cannot be further resolved (see Supplemental Fig. S15). (*F*) The representation of a bridged and an unbridged repeat in the de Bruijn graph approach, in the standard string graph approach and according to HINGE. The de Bruijn graph approach collapses the repeated segment, which allows a natural repeat resolution step if a bridging read is found. The representation in the string graph (if there is no read entirely contained in the repeat) is an hourglass-like motif. HINGE emulates the de Bruijn graph layout but in an overlap graph framework.

(Pevzner and Tang 2001; Mulyukov and Pevzner 2002; Butler et al. 2008; Peng et al. 2010).

In the context of third-generation long-read sequencing, however, standard de Bruijn graph approaches have not been as successful as they were in the case of short-read sequencing. Due to the high error rates associated with third-generation platforms, a large number of spurious k-mers is created, disrupting the structure of the de Bruijn graph. Recently, the concept of solid k-mers was proposed as a way to construct an "approximate" de Bruijn graph on a restricted set of reliable k-mers (Lin et al. 2016). However, since overlapping reads only share a handful of solid kmers, the resulting graph lacks the attractive features of de Bruijn graphs. In particular, the Eulerian structure is compromised and repeats are no longer properly collapsed into single paths. Overlapbased approaches, on the other hand, are more robust to read errors since they directly connect reads based on overlaps instead of first breaking them into k-mers. In fact, most available long-read assemblers (Chin et al. 2013, 2016; Berlin et al. 2015; Li 2016) are based on the so-called overlap-layout-consensus pipeline.

While de Bruijn graphs are Eulerian, overlap graphs are Hamiltonian; i.e., the underlying genome sequence corresponds to a cycle that traverses every node (read) in the graph. In addition to well-known computational challenges (Nagarajan and Pop 2009), the Hamiltonian paradigm does not yield a natural representation of repeat patterns, and the graph is typically riddled with unnecessary edges. In order to combat these issues, the string graph approach (Myers 1995, 2005) was proposed, originally for the Celera assembler (Myers et al. 2000; Myers 2016a) and later adopted by several assembly pipelines (Chin et al. 2013, 2016; Berlin et al. 2015; Li 2016). Built via a *transitive reduction* procedure, the string graph is an overlap graph where the unique, nonrepetitive parts of the genome correspond to simple, unbranched paths. However, long repeats—both bridged and unbridged—may result in undesirable graph motifs. In practice, only heuristics are used to combat these motifs, and building a maximally resolved overlap graph is challenging.

Results

We propose HINGE as a way to build an assembly graph where only the segments corresponding to unbridged repeats are collapsed. This objective, which we refer to as a maximally resolved assembly graph, is illustrated in Figure 1A–E. As depicted in Figure 1F, this goal is naturally achieved in a de Bruijn graph framework but not within an overlap graph-based framework due to the motifs created by long repeats. HINGE seeks to simultaneously attain the error resilience of overlap graph-based approaches and the appealing graph structure and optimal repeat resolution capability of de Bruijn graphs. Next, we briefly outline the main algorithmic innovations that allow HINGE to achieve this goal and present results on several data sets.

Algorithmic contributions

HINGE is an assembler that follows the overlap-layout-consensus paradigm. Its main algorithmic innovation lies in how it exploits the alignments obtained in the overlap phase in order to identify resolvable repeats and construct the graph layout in a repeat-aware fashion. Next, we describe the main ideas that go into the layout step. We defer a description of the overlap and consensus steps to the Methods section.

Repeat annotation and hinging reads

HINGE utilizes the alignment information obtained in the overlap step in order to equip some of the reads with *hinges*. Hinges are placed at the beginning and end of unbridged repeats and will ultimately lead to bifurcations on the graph, as illustrated in Figure 1F. The first step toward hinging the reads, as illustrated in Figure 4A below, is to find sharp gradients in the number of alignments on a read and annotate them as beginning or end of repeats. Next, we identify reads that bridge a repeat by finding reads that have both an annotation for the beginning of a repeat and an annotation for the end of the same repeat. Finally, we spread the information of which repeats are bridged to other reads through a procedure that we term the *Contagion algorithm* (see Methods; Fig. 5, below).

Hinge-aided greedy overlap graph construction

The Contagion algorithm allows HINGE to place exactly one inhinge and one out-hinge on the reads that originated from unbridged occurrences of a repeat. HINGE can then create a sparse overlap graph by using a hinge-aided greedy graph construction. In essence, we pick a best predecessor and a best successor for each read, as in the classical greedy algorithm (Tarhio and Ukkonen 1988) or in the best-overlap-graph approach (Miller et al. 2008). However, since our reads are hinged, we also allow a read's successor or predecessor to be the *interior* of another read, as long as the match starts on a hinge. When this occurs, a bifurcation is formed on the graph, corresponding to the beginning or the end of an unbridged repeat.

As illustrated in Figure 1F, this hinge-aided approach allows us to obtain the attractive properties of a de Bruijn graph layout but within the OLC framework. A comparison with the traditional greedy approach is provided in Supplemental Figure S2. We point out that for higher fold repeats, where a subset of the copies may be unbridged, a more careful handling of hinges is required, and that is achieved using a new procedure that we call *Poisoning*, described in the Methods section and in Figure 6 below.

Validation of hinge on data sets with ground truth

In Supplemental Figures S3–S6, we present validation results on simulated data sets. We created sequences with specific patterns of repeats and simulated long error-prone reads, using the DAZZ-DB simulator. We then verified that, when run on these data sets, HINGE produces a maximally resolved assembly graph. In Supplemental Figure S7, we validate the structural integrity of our assembly on an Oxford Nanopore R9 *Escherichia coli* data set. In Supplemental Figure S8, we validate the structural and sequence integrity of our assembly on a Pacific Biosciences (PacBio) *Saccharomyces cerevisiae* data set.

In Supplemental Table S2, we present validation results on *E. coli* data sets produced by PacBio and Oxford Nanopore sequencers. In both of these cases, HINGE produces a single circular contig and there is no misassembly. We also compare our assembly with the assembly produced by the NCTC pipeline (HGAP followed by Circlator) on 10 randomly selected data sets. We verify that the assemblies agree and have high identity scores in all cases.

Evaluation on the NCTC database

We evaluated HINGE on the 997 bacterial genomes of the NCTC 3000 database that were publicly available at the time of writing this manuscript (http://www.sanger.ac.uk/resources/downloads/bacteria/nctc/). The accession number for these data sets is provided in Supplemental Tables S1 and S3. Each of these data sets consists of PacBio SMRT long reads with coverage depths mainly in the range 30× to 80×. While the repeat complexity is relatively mild in bacterial genomes, we chose to evaluate HINGE on these data sets for two reasons: it allows us to carefully verify whether the HINGE assembly graphs satisfy our goal of maximal repeat resolution, and it allows us to run experiments on a large number of data sets, thus avoiding overfitting.

The current NCTC manual assembly pipeline uses the HGAP assembler (Chin et al. 2013) to produce a list of contigs, and Circlator (Hunt et al. 2015) to circularize contigs. The assembly graphs produced by HINGE with no parameter tuning for each of these data sets are available online (http://web.stanford.edu/~gkamath/NCTC/report.html) and in Supplemental Table S4, along with the contig statistics of the NCTC pipeline results and the assembly graph produced by Miniasm (Li 2016). We point out that other state-of-the-art assemblers, in particular, FALCON (Chin et al. 2016), have runtimes above one order of magnitude greater than HINGE (see Supplemental Fig. S11), making a comparison on the entire NCTC database computationally prohibitive.

For 822 of the 997 available data sets, HINGE produced a finished nonfragmented assembly graph, with additional isolated small plasmids in many cases. In 40 of these data sets, HINGE identifies unresolvable repeats, and the final graph admits distinct traversals (see Table 1). In order to compare our results with those obtained by the NCTC manual pipeline, we restricted our attention to those data sets for which NCTC reports the results of their assembly. As shown in Table 2, even without a circularization tool, HINGE obtains significantly more finished assemblies than the NCTC pipeline.

Analysis of HINGE assembly graphs

Among the cases where HINGE produces an assembly graph with multiple traversals, we find many examples where the intuitive layout of the graph produced by HINGE resembles the idealized cases in Figure 1A-E and allows one to visually assess the unresolvable repeat pattern in the genome. Next, we analyze three such cases in depth and compare the graph produced by HINGE with the contigs produced by the NCTC pipeline. We see that, by focusing on obtaining a maximally resolved assembly graph rather than large contig N50 values, HINGE prevents several misassemblies the NCTC pipeline incurred. In Supplemental Figure S9, we present nine additional such cases. In Supplemental Figure S10, we present several cases where HINGE resolves all repeats, producing a finished circular assembly, while the NCTC pipeline instead fragments the assembly. In addition, in Supplemental Figures S12-S14 we provide the same comparisons but with FALCON (Chin et al. 2016) instead of the manual NCTC pipeline.

In Figure 2A, we examine NCTC11022 (*E. coli*). In this example, the incorrect resolution of a 20-kbp unbridged repeat by the NCTC pipeline (see Supplemental Fig. S16) causes the circular chromosomal contig to lose a 780-kb segment, returned as a separate contig. By first collapsing this repeat and then resolving it due to the existence of a unique traversal of the graph, HINGE produces a single large chromosomal contig of length 5.1 Mbp. The nodes



Figure 2. Analysis of HINGE graphs on selected data sets. By identifying unbridged repeats, collapsing them, and then performing resolutions based on uniquely traversable loops, HINGE prevents misassemblies and produces a user-friendly, interpretable assembly graph. We color the graph nodes according to their corresponding position on the NCTC pipeline contigs. (A) On NCTC11022, HINGE identifies an unbridged repeat, which is later resolved. (B) On NCTC9024, HINGE identifies an unbridged triple repeat (with one inverted copy), which cannot be resolved due to the existence of three distinct traversals of the graph. (C) HINGE identifies an unbridged triple repeat. (D) HINGE identifies an unresolvable repeat shared by two small plasmids.

in the HINGE graph are colored according to the position the corresponding reads align to in the NCTC pipeline contigs.

On the NCTC9024 data set (*E. coli*) (Fig. 2B), the NCTC pipeline returned two long contigs, one of 4.3 Mbp and one of 0.9 Mbp. The HINGE graph emphasizes the existence of a triple repeat which, upon further inspection (see Supplemental Fig. S17), is seen to be of length 20 kbp, unbridged, and with one inverted copy. Even though this repeat is unbridged, both the NCTC pipeline and FALCON resolve one of its copies but in distinct ways. As we point out in Supplemental Figure S9, incorrect resolution of an inverted repeat can produce a false inversion of a long contig. In fact, the NCTC assembly and the FALCON assembly disagree on the orientation of the yellow-to-orange segment, and one of them must be creating an incorrect inversion of more than 1 Mbp (the orange-to-yellow segment). By collapsing the repeat, HINGE avoids a potential misassembly.

In Figure 2C, we consider NCTC9657 (*Klebsiella pneumoniae*). In this example, the NCTC pipeline returned seven unidentified contigs (three large ones), but HINGE returns a single large chromosomal connected component and three small plasmids. In

this case, HINGE produces a graph motif characteristic of an unbridged triple repeat, similar to Figure 1D. As shown by a coverage analysis in Supplemental Figure S18A, this is indeed a triple repeat, and contig 1 of the NCTC pipeline incorrectly resolves it, creating a misassembly. In addition, we examine the plasmids produced by the NCTC pipeline in Figure 2D and note that two of them share an unbridged repeat (see also Supplemental Fig. S18B). Therefore, there are two possible resolutions (two plasmids or a single, longer plasmid), and HINGE keeps them merged on the graph to retain this unresolvable ambiguity. In Supplemental Figure S12, we verify that the performance of FALCON (Chin et al. 2016) on the examples in Figure 2, A and C, is similar to that of the NCTC pipeline.

As illustrated by these examples, HINGE seeks to construct a user-friendly, informative, overlap graph as its main output, as opposed to most OLC assemblers, which employ assembly graphs in their inner workings (Chin et al. 2013, 2016; Berlin et al. 2015) but focus on outputting a list of contigs. To the best of our knowledge, Miniasm (Li 2016) is the only other assembler to produce a graph as the main assembly output. However, Miniasm is based on the string graph paradigm, which does not achieve the graph layout HINGE strives for, as we empirically observe (http://web.stanford. edu/~gkamath/NCTC/report.html).

Discussion

With HINGE, we introduce a new approach to constructing assembly graphs

in a repeat-aware fashion. While other state-of-the-art assemblers do attempt to identify bridging reads (sometimes referred to as spanning reads) and resolve the corresponding repeats, this is usually done as a post-processing step on the graph. HINGE, on the other hand, seeks to identify repeats and determine whether they should be collapsed on the graph prior to the actual construction. This way, HINGE avoids having to identify and correct graph motifs (such as the ones created by the string graph as shown in Fig. 1F) in a post-processing phase, which can be difficult due to spurious and missing edges caused by the high error rates of long-read sequencing technologies and by chimeric reads.

In order to reliably achieve this repeat-aware graph layout, several new conceptual ideas were introduced in HINGE. First, a repeat annotation step is responsible for identifying the beginning and end of repeats and which reads bridge some repeat. However, this type of local information is not sufficient for the construction of a maximally resolved assembly graph. Therefore, this information must be spread to other reads, which is accomplished with our Contagion algorithm.

	Coverage ≥40×	All coverages
Number of NCTC data sets HINGE finished circular assembly (single traversal) HINGE finished assembly (lacking circularization) HINGE finished circular assembly (multiple traversals)	$\left.\begin{array}{c} 816\\ 631\\ 60\\ 38\end{array}\right\} 691 \\ 729\\ 38\end{array}\right\}$	$ \begin{array}{c} 997 \\ 690 \\ 92 \\ 40 \end{array} \right\}782 $ 822

Given the output graph of HINGE we classify the assembly into four categories. A *finished circular assembly* corresponds to a case where all nodes (small plasmids excepted) lie on a single circle. A *finished circular assembly with multiple traversals* corresponds to a graph where all nodes can be visited by a circular path, but there is more than one such path. We point out that we classify such an output as finished because such a graph can be seen as simultaneously capturing a few (usually two) assemblies, all of which would be considered finished according to the previous rule. A finished assembly is said to *lack circularization* if a single noncircular path can traverse all nodes on the graph (small plasmids excepted). If the graph produced by HINGE does not fall into the previous three categories, we classify it as a misassembly/fragmented assembly. As reliable hinge placement requires a reasonable coverage depth, we also considered restricting our attention to the data sets with average coverage depth above 40. We note that Miniasm needs a circularization tool to circularize assemblies, and hence, we report a Miniasm assembly as finished if it has only one contig longer than 200 kbp. The graph produced by HINGE and Miniasm for all these cases can be found at http://web.stanford.edu/~gkamath/NCTC/report.html and in Supplemental Table S4, along with the corresponding classification. As can be seen on this report, the rule for determing when a Miniasm assembly is finished is often quite lenient.

Once the bridging information is known globally, HINGE utilizes a hinge-aided greedy construction of the graph. This is also different from most state-of-the-art long-read assemblers, which rely on the string graph paradigm. Our approach bears similarities with the best overlap graph approach in its goal of constructing a sparse overlap graph but takes advantage of hinges as a way to achieve this goal with maximal repeat resolution. Finally, the sparse nature of the constructed graphs allows HINGE to identify loops that admit a single traversal and can thus be resolved. The conceptual contributions of HINGE are discussed in more detail in the Methods section.

As an OLC assembler, in order to produce high quality assemblies, HINGE relies on good Overlapping and Consensus modules. In its current implementation, HINGE was designed to work with the output of DALIGNER (Myers 2014), and the consensus is performed using a variant of the consensus module of FALCON (Chin et al. 2016) together with a straightforward majority-vote finishing step. These choices are not essential to the workings of our pipeline. Therefore, integrating HINGE with other overlapping tools such as MHAP or Minimap can be done if different levels of alignment sensitivity or memory usage are required. Similarly, different consensus and polishing modules such as Quiver (Chin et al. 2013) and Racon (Vaser et al. 2017) can be used, according to the desired point in the accuracy-computation tradeoff.

Through a novel approach to repeat resolution and graph representation, HINGE brings a fresh perspective to the assembly problem. By focusing on the construction of a maximally resolved assembly graph in a user-friendly fashion, HINGE is well aligned with the recent push for the standardization of graph references, as opposed to the traditional contig representation. The HINGE graph is a natural representation of a set of possible assemblies and is amenable to further repeat resolution, which can be attempted using additional long-range information such as pairedend reads, Hi-C reads, or by leveraging biological insight. Finally, we point out that, while the repeat complexity is relatively mild in the bacterial genomes we consider (as evidenced by the large number of finished assemblies), it is much more severe in higher organisms (Koren et al. 2013). This highlights the importance of the careful treatment of repeats carried out by HINGE and the value of the proposed method to genome assembly.

One important aspect regarding the notion of maximal repeat resolution is that it assumes that long contiguous matches identified in the read alignment step must correspond either to the same segment on the genome or to repeats whose copies are similar enough that they should be merged in the graph. However, there may still be a small level of divergence between these copies that is below the sequencing error rates and cannot be detected by the aligner. In principle, this divergence may allow a final "phasing" or "unzipping" step, similar to what is used in FALCON-Unzip (Chin et al. 2016), to resolve these repeats. Utilizing these small levels of divergence to phase or to score the different traversals of a repeat according to their likelihood is a future direction for improvement of the HINGE pipeline.

Methods

The HINGE assembly pipeline is an OLC pipeline designed to assemble long reads. The overall workflow is depicted in Figure 3

Table 2. Finished assemblies on NCTC data sets where NCTC manual pipeline results are reported

	Coverage ≥40×	All coverages
Number of NCTC data sets	688	834
NCTC manual pipeline finished assemblies	517	592
Miniasm finished assembly (not circularized)	513	592
HINGE finished circular assembly (single traversal)	531) ₅₈₃	583 } 660
HINGE finished assembly (lacking circularization)	₅₂ J ⁵⁰⁵	77 J 000
HINGE finished circular assembly (multiple traversals)	33	33

In this table, we restrict the data sets considered in Table 1 to only those for which NCTC reports a result for comparison. The finished assemblies for the NCTC manual pipeline correspond to the cases where they report one chromosomal contig or two chromosomal contigs (since species such as *Vibrio fluvialis* and *Ochrobactrum anthropi* are known to have two chromosomes). We point out that, while a circularization tool (Circlator) is used in the NCTC pipeline, we do not have a circularization finishing step and only report the output of HINGE using default configurations.



Figure 3. HINGE pipeline. (*A*) The input to the HINGE pipeline is a set of long error-prone reads. (*B*) Chimeric reads are detected through their pile-o-grams and are discarded. (*C*) The beginning/end of repeats are annotated on the reads. This is done by detecting a sharp increase/decrease in the number of alignments on a read. The repeat annotations are also identified as bridged or unbridged. (*D*) Maximal reads (i.e., reads that are not contained in other reads) are selected and fed to the Contagion algorithm, which is responsible for spreading the information about which repeats are bridged to all the reads, allowing us to place exactly one in-hinge and one out-hinge on the reads that originated from unbridged occurrences of a repeat. (*E*) The set of maximal reads (some of which are not hinged) is the input to the hinge-aided greedy assembly. (*F*) After obtaining the reads onto the resulting overlap graph, we use standard consensus methods to generate contigs.

and is explained in detail in this section. As the default parameters and auxiliary tools were selected to optimize the pipeline for PacBio reads, we focus the discussion on this setting.

Read database and alignment

We use DAZZ_DB (Myers 2016b) to maintain a database of the PacBio reads. We use DALIGNER (Myers 2014) to obtain pairwise alignments between all reads. We point out that HINGE does not heavily rely on specifics of the DALIGNER output and can be adapted to work with other aligners as well.

No initial error-correction step

Unlike most available long-read assembly pipelines, HINGE bypasses an initial error-correction step. To the best of our knowledge, Miniasm (Li 2016) is the only other OLC assembler that dispenses with this step. ABruijn (Lin et al. 2016) also has no error-correction step, though it is not based on the OLC paradigm. The idea of error-correction-free assembly was also utilized in Lien et al. (2016) and Tørresen et al. (2017). The fact that longread aligners like DALIGNER (Myers 2014) can obtain pairwise alignments at error rates around 15% allows us to use this approach and defer the error correction to the final consensus step.

Chimeric read filter

Chimeric reads are the result of a sequencing error and are usually made up of multiple segments that originate from different

parts of the genome. If not properly handled, these reads create misassemblies, and different techniques have been put forward to detect chimeras (Miller et al. 2008; Li 2016). HINGE's chimera filter unit is the first place in the pipeline where the visualization provided by pile-o-grams (Supplemental Figs. S19, S20) is useful. We mark a read segment as chimeric if the set of reads aligned to it undergoes an abrupt change. On the pile-o-gram, as shown in Figure 3B, one sees a clear discontinuity in the set of alignments (blue segments) of a read. We also mark a read segment as chimeric if the number of matches goes below a fixed threshold. For each read, we keep the longest segment without any chimeric segments. If this segment is shorter than a threshold, we discard the read completely.

Repeat annotation

One of the main distinctive features of HINGE is a pre-assembly step responsible for annotating the beginning and the end of repeats on the reads. These repeat annotations will later be used for placing hinges on the reads, which, in turn, will be instrumental in the graph layout step. The repeat annotation is done by detecting the start/end of a large number of matches on a read. On the pile-o-gram (Supplemental Figs. S19, S20), this visually corresponds to a large pile of matches starting/ending at the same point, as

shown in Figure 3C. We note that relying on coverage gradients rather than coverage itself makes HINGE immune to coverage fluctuations.

We then verify whether the repeat annotation corresponds to a repeat that is bridged by that read. Intuitively, one could attempt to do this by identifying both a sharp increase and a sharp decrease in the number of matches on a read. However, as it turns out, such an approach can fail in the presence of more complex repeat patterns such as a repeat within a longer repeat (see Supplemental Fig. S19E for an illustration). Therefore, a more careful processing of the matches on a read is needed to identify the bridging condition. HINGE determines the bridging condition by checking whether most of the matches starting on a repeat annotation also end on a repeat annotation. If that is the case, the repeat is assumed to be bridged, and the annotation is flagged as such (red annotations in Fig. 3C). Thus, at the end of this step, we have repeat annotations on all reads, and these annotations are labeled as bridged/unbridged according to the local information provided by the reads' alignments. The next step, the Contagion algorithm, is applied to this set of annotated reads after we filter out reads that are fully contained in other reads (keeping only maximal reads).

The Contagion algorithm

Notice that this local information about the bridging of repeats may be misleading. For example, the pile-o-gram of read u in Figure 4A may suggest that u lies partially on an unbridged repeat. However, that repeat might still be bridged by a different read, as in



Figure 4. (*A*) Sharp changes in the number of alignments give rise to repeat annotations on each read. If a read is verified to bridge a repeated, as in the case of read v, the corresponding read annotations are marked as such (shown as red nodes). (*B*) The Contagion graph is formed by having all repeat annotations as nodes, and using edges to mark annotations that correspond to the beginning (or end) of the same repeat. As illustrated here for NCTC11022, connected components with no bridged repeat annotations will give rise to hinged reads, which leads to bifurcations on the graph. The repeats corresponding to other connected components stay resolved in the graph.

the case of read v in Figure 4A. Therefore, HINGE proceeds to "spread" the local bridging information of each read to other reads using the Contagion algorithm. At a high level, this algorithm can be thought of as constructing a Contagion graph (see Fig. 4B), with nodes being the repeat annotations and edges between repeat annotations that correspond to the beginning (or end) of the same repeat (possibly from different copies of the same repeat). Annotations corresponding to the beginning/end of the same repeat are identified based on alignments: if two reads have an annotation corresponding to the beginning (respectively, end) of a repeat and have matching segments after (respectively, before) the annotation, the two annotations are connected in the graph. The edge points in the direction of the read that extends the most into the repeat. Moreover, repeat annotations that have been identified as the beginning/end of a bridged repeat are marked as such (red nodes in Fig. 4B).

As illustrated in Figure 4B, this graph has two connected components for each repeat (the yellow and pink components correspond to the beginning and end of an unbridged repeat). In this graph, repeat annotations corresponding to bridged repeats are thought of as "infected" (shown as red nodes) and can spread the "bridging condition" to the other repeat annotations in the same connected component. If a connected component does not contain any infected repeat annotation, it corresponds to the beginning/end of an unbridged repeat and will eventually lead to a bifurcation on the graph (see hinge-aided greedy assembly algorithm), as shown in Figure 4B.

The Contagion algorithm processes the Contagion graph to kill repeat annotations that will not be useful in the overlap graph construction. In particular, this infecting and killing process performs two tasks: (1) repeat annotations corresponding to bridged repeats should cause other annotations corresponding to the same repeat to also be marked as bridged and ultimately killed; (2) if two repeat annotations correspond to the beginning (or end) of the same repeat, the one extending the most into the repeat should be kept, while the other one should be killed. This global processing of the repeat annotations and bridging condition is important so that ultimately we only place one in-hinge and one out-hinge for each unbridged repeat (on the sink node of the corresponding connected component).

In more detail, the Contagion algorithm comprises three steps. In the first step, we remove all annotations whose connected component on the Contagion graph is small. For instance, the gray-colored nodes in Figure 4B correspond to small connected components that are deleted. Typically, these small components are the result of imprecise placement of repeat annotations on reads, which then lead to them not matching other repeat annotations that correspond to the beginning/end of the same repeat. Hence, deleting these small components prevents us from creating multiple hinges corresponding to the beginning/end of the same repeat.

In the next step, we look for pairs of repeat annotations connected by an edge in the Contagion graph (i.e., corresponding to the beginning/end of the same repeat) and such that the corresponding reads have an overlap. (Notice that by an overlap, we mean a match between the suffix of a read and the prefix of another read. If the match instead occurs in the interior of at least one of the reads, we refer to it as an internal match.) For every such pair, we kill the annotation on the read that extends the least into the repeat. For an illustration of this step, consider the two unbridged repeats in Figure 5A. The reads covering the start of each repeat (u_1, u_2) u_2, u_3) have a start-repeat annotation at the start of the repeat. The reads covering the end of each repeat (v_1, v_2, v_3) have an end-repeat annotation at the end of the repeat. As shown in Figure 5B, the start-repeat annotation on u_2 is killed by the start-repeat annotation on u_1 because u_2 and u_1 have an overlap and u_2 extends more into the repeat. Similarly, the end-repeat annotation at v_2 is killed due to the overlap with v_1 . At the end of this step, we have that exactly one read covering each copy of an unbridged



Figure 5. The Contagion algorithm. (*A*) Two unbridged repeats are shown as orange segments. (*B*) The Contagion algorithm first kills the start-repeat annotation at u_2 (due to its overlap with u_1) and the end-repeat annotation at v_2 (due to its overlap with v_1). (C) The Contagion algorithm then kills the start-repeat annotation on u_3 (due to its internal match with u_1) and the end-repeat annotation on v_3 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_1) and the end-repeat annotation on v_1 (due to its internal match with u_2) and u_3 to choose the in-hinge at u_1 as their successor. Similarly, v_1 and v_2 pick the match starting at the out-hinge on v_3 as their predecessor match.

repeat has a start-repeat annotation on it (and exactly one read has an end-repeat annotation on it for each copy of the repeat). In addition, we point out that when a read has its repeat annotation killed by an annotation from a bridged repeat, we mark this annotation as "poisoned." The reason for the term is that a poisoned read would be "deadly" for a standard greedy assembly algorithm, as it would lead to a misassembly.

The third step of the Contagion algorithm is similar to the second step, but instead of looking for matching annotations whose reads have an overlap, we look for matching annotations whose reads have an internal match. For every such pair of annotations, we keep the one on the read that extends the most into the repeat, and kill the other one. As illustrated in Figure 5C, this causes the start-repeat annotation on u_3 to be killed by the start-repeat annotation on u_1 and the end-repeat annotation on v_1 to be killed by the end-repeat annotation on v_3 . At the end of this step, we have one in-hinge on u_1 and one out-hinge on v_3 . We point out that, in this step, we only consider nonpoisoned reads.

Finally, all surviving annotations for the start of unbridged repeats are marked as in-hinges and the annotations for the end of unbridged repeats are marked as out-hinges. One can formally show that under the assumption that no significant alignment is missed in the initial Overlap step, the Contagion algorithm will place exactly one in-hinge and one out-hinge among the reads that originated from the set of unbridged occurrences of a repeat and no hinge on the reads from the bridged occurrences of a repeat.

Hinge-aided greedy assembly algorithm

A key distinction of HINGE's approach to assembly lies in its graph layout step. Many OLC assemblers adopt the string graph paradigm (Myers

1995, 2005), which often produces assembly graphs that are unnecessarily dense. HINGE replaces the string graph algorithm with a variant of the greedy algorithm. This follows a recent line of work that found that variants of the greedy algorithm (such as the best-overlap-graph [BOG] algorithm [Miller et al. 2008], "not-so-greedy" algorithm [Shomorony et al. 2016b], and the greedy merging algorithm [Shomorony et al. 2016a]) can produce a sparse overlap graph without misassemblies.

Notice that, at the end of the Contagion algorithm, we only have one in-hinge and one out-hinge for each unbridged repeat. In the graph layout step, we employ a variation of the greedy algorithm that utilizes the hinge information. Each read picks its left extension to be its longest prefix match and its right extension to be the longest suffix match. However, unlike in the classical greedy algorithm, we do not restrict our search to overlaps. In addition to (prefix-suffix) overlaps, we also consider internal matches. Hence, a read is allowed to find its successor/predecessor match to be an internal segment of another read

as long as the match starts on a hinge. An illustration of how internal matches are helpful in producing the correct graph layout and a comparison with the classical greedy algorithm is illustrated in Supplemental Figure S2.

The role of poisoned reads

Another important aspect of the hinge-aided layout step is how the read poisoning information is used. As mentioned above, in the Contagion algorithm, whenever a read has its start/end repeat annotation killed by another overlapping read, we label it as poisoned. During the hinge-aided greedy algorithm, reads are prevented from picking a poisoned read as their predecessor/ successor, guaranteeing that the two copies (or occurrences) of a bridged repeat remain separate.

This process is illustrated in Figure 6. In the scenario shown in Figure 6A, read u_1 is initially given a start-repeat annotation. However, this start-repeat annotation is removed by the





Contagion algorithm, as the repeat is bridged by read w. In this case, we keep a poisoned annotation on read u_1 . When a read has a poisoned start-repeat annotation, it cannot be chosen as a predecessor of another read if the match starts after the start-repeat annotation. As shown in Figure 6B, according to a non-hinge-aided greedy assembly algorithm, v_2 would choose u_1 as its predecessor. However, as the match on u_1 starts after the poisoned start-repeat annotation, we do not allow v_2 to choose u_1 as a predecessor. Instead, we look for the next best option, which in this case is u_2 . This prevents a misassembly. The poisoning of end-repeat annotations works in an analogous way.

In addition, we point out that the concept of poisoning is what allows the proper collapsing of the unbridged copies of repeats with three or more copies. Notice that during the third stage of the Contagion algorithm, we only consider nonpoisoned reads. Therefore, we only deal with reads coming from unbridged copies of the repeat. As a result, the set of all unbridged occurrences of a repeat induces exactly one in-hinge and one out-hinge. On the other hand, all reads from the bridged copies are poisoned and receive no hinge.

Repeat resolution

Another new ingredient introduced by HINGE is the use of global information to resolve repeats. Once constructed, the graph allows us to identify certain repeats that, although unbridged, can still be resolved based on the graph layout. As illustrated in Figure 3A and Supplemental Figure S15A, when a repeat loop allows only one possible traversal, the loop can be untangled. We point out that the sparse and Eulerian-like nature of the graph produced by the hinge-aided greedy algorithm is important to allow this repeat resolution to be done in an automated fashion. We also point out that the loop resolution step is based on a parsimony principle, but it could be potentially incorrect if the loop corresponds to a plasmid, to a separate chromosome, or to the genome of a different species present in the sample. The parameter MAX_PLASMID_LENGTH sets the maximum length of a loop that should be considered a potential plasmid. HINGE will only resolve loops longer than MAX_PLASMID_LENGTH, and this behavior can be optionally turned off by setting MAX_PLASMID_LENGTH to a number longer than the genome length.

Handling read orientation and double-strandedness

Since the orientation of the reads is unknown, as is typical in all assembly pipelines, one must consider each read and its reverse complement. Hence, for each read, we in fact create two nodes in the graph, and the constructed graph is symmetric. At the end of the graph construction, for visualization purposes, we overlay each node and its reverse complement.

Consensus

In order to generate consensus sequences for the resulting graph contigs, we first create a draft assembly by simply concatenating sections of the error-prone reads corresponding to unbranched paths on the graph. We then consider the alignment of all the original reads onto these draft contigs and utilize a simple majoritybased consensus to clean up these draft sequences. We reuse some code from FALCON (Chin et al. 2016) to perform this task. The result is output as a GFA file. We point out that the final contig sequences can be optionally run through Quiver (Chin et al. 2013) to further polish the assembly.

Graph visualization

All assembly graphs produced by HINGE were visualized using Gephi (Bastian et al. 2009).

Software availability

The HINGE assembler is available online at https://github.com/ HingeAssembler/HINGE and in the Supplemental Source Code. The analyses presented in Figure 2 can be reproduced in https ://github.com/HingeAssembler/HINGE-analyses.

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Author contributions: G.M.K. and I.S. designed the algorithm. F.X. implemented a testbed to test and experiment with assembly algorithms. G.M.K., I.S., and F.X. implemented the HINGE algorithm, ran it on the data set, and visualized and interpreted the results. T.A.C. and D.N.T. supervised the project. All authors wrote the paper.

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