DENTIST – using long reads to close assembly gaps at high accuracy

Arne Ludwig 1,2, Martin Pippel 1,2, Gene Myers 1,2, Michael Hiller 1,2,3,4,5,6*

1 Max Planck Institute of Molecular Cell Biology and Genetics, Pfotenhauerstr. 108, 01307 Dresden, Germany
2 Center for Systems Biology Dresden, Pfotenhauerstr. 108, 01307 Dresden, Germany
3 Max Planck Institute for the Physics of Complex Systems, Nöthnitzer Str. 38, 01187 Dresden, Germany
4 LOEWE Centre for Translational Biodiversity Genomics, Senckenberganlage 25, 60325 Frankfurt, Germany
5 Senckenberg Research Institute, Senckenberganlage 25, 60325 Frankfurt, Germany
6 Goethe-University, Faculty of Biosciences, Max-von-Laue-Str. 9, 60438 Frankfurt, Germany

*To whom correspondence should be addressed:
Michael Hiller
LOEWE Centre for Translational Biodiversity Genomics, Senckenberganlage 25, 60325 Frankfurt, Germany
Tel: +49 69 7542-1398
E-Mail: Michael.Hiller@senckenberg.de

Running title: DENTIST – accurately closing assembly gaps

Keywords: genome assembly, long sequencing reads, assembly gaps, scaffolding
Abstract
Long sequencing reads allow increasing contiguity and completeness of fragmented, short-read based genome assemblies by closing assembly gaps, ideally at high accuracy. While several gap closing methods have been developed, these methods often close an assembly gap with sequence that does not accurately represent the true sequence.
Here, we developed DENTIST, a sensitive, highly-accurate and automated pipeline method to close gaps in short read assemblies with long reads. DENTIST comprehensively determines repetitive assembly regions to identify reliable and unambiguous alignments of long reads to the right loci, integrates a consensus sequence computation step to obtain a high base accuracy for the inserted sequence, and validates the accuracy of closed gaps. Unlike previous benchmarks, we generated test assemblies that have gaps at the exact positions where real short-read assemblies have gaps. Generating such realistic benchmarks for Drosophila (134 Mb genome), Arabidopsis (119 Mb), hummingbird (1 Gb) and human (3 Gb) and using simulated or real PacBio reads, we show that DENTIST consistently achieves a substantially higher accuracy compared to previous methods, while having a similar sensitivity. As another distinguishing feature, DENTIST can accurately scaffold the given contigs with long reads in addition to closing gaps, extending its application range to contig-only assemblies.
In summary, DENTIST provides an accurate approach to improve the contiguity and completeness of fragmented assemblies with long reads. DENTIST’s source code including a Snakemake workflow and Docker container is available at https://github.com/a-ludi/dentist. All test assemblies as a resource for future benchmarking are at https://bds.mpi-cbg.de/hillerlab/DENTIST/.
Introduction

The quality of a genome assembly has an important impact on the quality of any downstream genomic analysis. High contiguity, completeness and accuracy of genome assemblies is fundamental to (i) comprehensively annotating genes, their promoters and other functional genomic elements, (ii) understanding genomic repeat content and organization, (iii) performing phylogenomic analysis, (iv) linking phenotypic differences to genomic differences, (v) mapping transcriptomic or epigenetic read data to an assembly, and ultimately using genomes for evolutionary and biomedical studies [1-3]. However, assembly of complex genomes is a challenging task, because sequencing reads are much shorter than chromosomes and large parts of eukaryotic genomes consist of repetitive regions [4]. Facilitated by advances in short read sequencing that drastically increased throughput and sharply decreased costs, genomes of numerous species have been sequenced with short read technologies such as Illumina instruments [5, 6]. However, such reads are too short to span many repetitive regions, resulting in rather fragmented assemblies with short contigs (contiguous sequences) despite high read coverage. Scaffolding methods such as mate pair sequencing or chromosome conformation capture (Hi-C) read pairs can order and orient contigs into long, sometimes chromosome-scale scaffolds. Nevertheless, due to a large number of assembly gaps, i.e. unknown genomic sequence between neighboring contigs, such assemblies remain inherently incomplete.

Pacific Biosciences (PacBio) or Oxford Nanopore Technologies instruments enable long sequencing reads, which can cover many kilobases and sometimes exceed one megabase in length. Long reads provide an effective and cost-efficient means to close gaps in fragmented assemblies, since these reads are often longer than genomic repeats, allowing one to bridge between neighboring contigs thus closing an assembly gap [7]. To close gaps in existing, more fragmented assemblies with long reads, several methods have been developed in the past, exemplified by PBJelly [8], FinisherSC [9], PacBio GenomicConsensus (PacBio GC; comprising the Arrow and Quiver methods; https://github.com/PacificBiosciences/GenomicConsensus), LR_gapcloser [10] and others [11-13]. These methods align a set of given long reads to an input assembly, determine which reads span assembly gaps and close these gaps with the new sequence. However, as we demonstrate below, even for shorter and less repeat-rich genomes, these tools lack high accuracy when closing assembly gaps. Thus, while applying these tools improves assembly contiguity, they compromise the quality of the resulting assembly, which can impair downstream analyses.

Here, we developed DENTIST, a new, reliable and sensitive gap closing method that unlike previous methods achieves a high level of accuracy. Using various simulated and real data sets for genomes ranging from small (Drosophila) to large and complex (human), and using realistic assembly gap loci, we demonstrate that DENTIST consistently achieves the highest accuracy compared to other state-of-the-art approaches, while having a similar or better
runtime and memory consumption. Furthermore, DENTIST is able to use long reads to scaffold the given contigs, extending its application to assemblies that consist only of contigs.

Results
Overview of DENTIST
DENTIST implements a full gap closing pipeline (Figure 1) and was developed with the main consideration of closing assembly gaps at a very high accuracy. Conceptually, given an assembly and a set of long reads, gap closing starts by aligning the reads to the assembly to identify those that reach into or span assembly gaps. The reads are then used to infer the DNA sequence that should be used to fill these gaps.

DENTIST differs from previous approaches in the following key aspects. First, a key step in accurately closing gaps is to align the long reads to the right loci in the input assembly. This is not a trivial task, since genomic repeats can lead to ambiguous read alignments and potentially assigning reads to the wrong genomic locus. Therefore, DENTIST integrates four approaches to identify repetitive regions in both the input assembly and the input long reads, and explicitly uses this repeat mask to identify reliable read alignments. Many of the remaining alignment ambiguities and conflicts are resolved using a scaffolding graph. Second, as long reads generally have a high base error rate, it is important to determine an accurate consensus sequence from the long reads that span or overlap an assembly gap before closing the gap. DENTIST employs a state-of-the-art reference-based consensus caller to generate high-quality consensus sequence for each closed assembly gap, maintaining a high base accuracy in the final assembly. Third, aiming at a high accuracy, DENTIST validates both the consensus sequence as well as the closed gap by aligning the input reads again to the gap-closed assembly, and does not perform questionable gap closures in favor of a correct result. Fourth, unlike other tools, DENTIST can also use long reads to scaffold the given contigs, enabling its application to contig-only assemblies.

Test procedure
To assess the performance of DENTIST and compare it to existing methods, we followed previous strategies to generate a “ground truth scenario” where assembly gaps were artificially-inserted into a high quality assembly and long reads were used to close the created assembly gaps. This general strategy allows one to compare sensitivity (number of closed gaps, increase in assembly contiguity) and, since the real gap sequence is known, the accuracy of the inserted sequence.

Unlike previous comparisons, we devised a ground truth scenario, where the assembly gap sizes and loci are as realistic as possible, which is important to assess real life performance. Previous comparisons introduced assembly gaps at random locations [8] or replaced repeats by assembly gaps of the same length [10]. By excising repeats, the latter approach results in
gap flanks having often unique, non-repetitive sequence that make accurate read alignment easier. Consequently, both approaches do not reflect the complexity of assembly gap sizes and locations in reality, and the fact that contig ends often reach into the unbridgeable repeat (as opposed to ending right before it). To overcome this, we devised a procedure that uses two real assemblies of the same species. A high quality assembly constitutes the “ground truth” and a fragmented short read assembly is used to obtain realistic assembly gap locations. Specifically, we aligned the fragmented assembly to the ground truth assembly and introduced gaps into the ground truth assembly at the exact position and size where gaps occurred in the short read assembly (Figure 2). Then we either used long reads that were sampled (simulated) from the ground truth assembly or real PacBio reads (Supplementary Table 1) to close assembly gaps and evaluate sensitivity and correctness.

In the following comparison, we applied DENTIST and three state-of-the-art methods, PBJelly [8], FinisherSC [9], PacBio GenomicConsensus (https://github.com/PacificBiosciences/GenomicConsensus), and LR_gapcloser [10] to four species with various genome sizes and repeat content, representing the insect, plant and vertebrate lineage: the fruit fly Drosophila melanogaster with a 134 Mb assembly where ~10% of the ground truth assembly was covered by DENTIST’s repeat masks, the thale cress Arabidopsis thaliana (119 Mb, ~11% repeats), the hummingbird Calypte anna (1 GB, ~2% repeats), and human (3 Gb, ~20% repeats) (Supplementary Table 1). We were only able to test FinisherSC on the smallest genome (Drosophila), as this method failed on Arabidopsis with simulated read, did not finish after two days on Arabidopsis with real PacBio reads and required more than 1 TB of RAM for larger genomes.

Simulated Read Data
We first simulated PacBio reads from each ground truth assembly with a typical PacBio error profile and an average raw read length of 27 kb (Supplementary Table 1). For Drosophila, Arabidopsis and hummingbird, the gap sequence inserted by DENTIST is on average ≥99.8% identical to ground truth sequence (Table 1, Figure 3A). In comparison, the inserted sequences by PBJelly were 88.61%–95.17% identical and the inserted sequences by LR_gapcloser had only ~87.7% average identity to the ground truth sequence. FinisherSC, which we could only run on the Drosophila dataset, had an identity of 81.53% (Table 1). For the 3 Gb human genome, the average sequence identity for DENTIST was 98.70%, whereas PBJelly (89.25%) and LR_gapcloser (88.01%) achieved lower average identity values. In addition to a high average identity, DENTIST closed 46.1% (human) to 91.1% (Drosophila) of the gaps with a 100% accurate sequence, while other methods at best closed 3.58% (PBJelly, 48 of 1,340 gaps for Drosophila) at 100% accuracy.

To account for the different sizes of assembly gaps, we also calculated the average identity weighted by the size of the gap. This metric showed highly similar values (Table 1), with the exception of PBJelly where the weighted average identity increased by 0.94–4.80%, indicating that larger gaps tended to be closed with a higher accuracy (discussed below). Nevertheless,
the maximum weighted average of PBJelly (96.11% for Arabidopsis) is lower than the accuracy of DENTIST. In summary, consistently for all assemblies, DENTIST achieves the highest accuracy.

Comparing sensitivity, we found that PBJelly closed the most gaps for Drosophila, LR_gapcloser closed most gaps for Arabidopsis, but both methods and DENTIST closed ~90% or more of the gaps in these two assemblies (Table 1, Figure 3A). For the two larger genomes, DENTIST closed the most gaps (hummingbird 98.7% of the gaps, human 78.9%) than other methods. To measure the increase in assembly contiguity, we computed the contig NG50 value, using the number of real bases (A,C,G,T) in the ground truth assembly as a fixed reference. This allows us also to compute the maximally possible contig NG50 that can be achieved by closing all introduced gaps in the ground truth assembly. It should be noted that contig NG50 increase is not a perfect measure of sensitivity, as the NG50 increase heavily depends on the position of closed gaps and reflects only indirectly the number of closed gaps. For example, closing a single gap between two large contigs could result in a larger NG50 increase than closing several gaps between several small contigs. Indeed, this effect is also visible in our comparisons, exemplified by the hummingbird assembly where LR_gapcloser achieved a higher contig NG50 (14.6 Mb vs. 12.8 Mb) despite DENTIST closing 3,118 more gaps. Despite these caveats, comparing contig NG50 increase, we found LR_gapcloser achieved the highest contig NG50 for Drosophila and hummingbird, PBJelly achieved the highest contig NG50 for Arabidopsis, and DENTIST achieved the highest contig NG50 for human. For all four species, DENTIST substantially increased the contig NG50 between 5-fold (Arabidopsis) and 259-fold (hummingbird). We note that PBJelly outputs an Arabidopsis assembly with a significantly higher contig NG50 value than what is achievable by correctly closing all introduced gaps (8.3 vs. 7.1 Mb), which indicates that at least some gaps were “overfilled”. Since NG50 reduces contiguity to a single number, we also plotted which percent of the assembly is contained in contigs exceeding a certain size (Figure 3C). These NG(x) plots showed that DENTIST, PBJelly and LR_gapcloser achieve similar improvements in contiguity for Drosophila and Arabidopsis, and that DENTIST and LR_gapcloser achieve a higher contiguity for hummingbird and human. In summary, our comparisons using simulated reads show that the sensitivity of DENTIST is comparable and sometimes better than other methods and that DENTIST excels in a very high accuracy.

Real PacBio long read data
Since simulated reads cannot capture the full diversity of issues that are present in real PacBio reads (chimeras, low quality regions, missed adapters) and further do not encompass the whole genome but were only sampled from the ground truth contigs, we next evaluated the performance of DENTIST and other gap closing methods on the same assemblies but using real PacBio reads (Table 1). We added the PacBio GC method to the tests, which requires additional sequencing metrics (such as pulse widths and inter-pulse durations) as input and can therefore only be applied to real read data. We note that in these tests, it will be harder to achieve a very high accuracy as real reads also contain SNPs or larger haplotype variation
and the ground truth assembly may contain base or assembly errors with respect to the real reads. Nevertheless, the accuracy of DENTIST is still very high with an average identity between inserted and ground truth sequence of 99.95% for *Drosophila*, 99.35% for *Arabidopsis*, 98.41% for the hummingbird and 96.66% for human (Table 1, Figure 3B). All other methods have a lower accuracy. A few examples of closed gaps are illustrated in Supplementary Figure 1.

For human, DENTIST’s average identity weighted by gap size is higher than the unweighted identity (98.02% vs. 96.66%), indicating that some of the smaller gaps were closed less accurately than larger gaps, which we confirmed by plotting identity vs. gap size (Figure 4). Manual inspection showed that many of these cases comprise gaps <50 bp consisting of simple repeats such as homopolymer runs, for which long reads have a higher error rate [14], making it more difficult to compute an accurate consensus. Since very few incorrect bases in the consensus of a small gap have a large effect on the average identity (e.g. 19 of 20 correct bases is an identity of only 95%), the weighted average identity provides a better measure of overall sequence accuracy. Furthermore, few base errors in a small gap are not a serious problem, since small gaps can easily be “polished” with Illumina reads in a downstream step (discussed below). A consistently higher weighted average identity was also observed for PBJelly (Table 1, Figure 4B).

Compared to DENTIST, the average identity (weighted and unweighted) of the sequence inserted by all other methods was consistently lower by at least a few percent. Furthermore, in contrast to DENTIST, other methods generate lower base identities not only for short gaps but especially for long gaps, and they rarely close long gaps with a very high accuracy (Figure 4).

In terms of sensitivity, LR_gapcloser and PacBio GC closed the most gaps and LR_gapcloser achieved the highest contig NG50 increases (Table 1, Figure 3D). DENTIST closed 74% (Human) to 94% (*Drosophila* and hummingbird) of the gaps, increasing contig NG50 by 3.3 times (*Arabidopsis*) to 33.0 times (hummingbird). Thus, consistent with the simulated read data, DENTIST has a reasonable sensitivity and the highest accuracy also when using real reads.

**Runtime and Memory Consumption**

While a high accuracy is certainly the most important objective, speed and memory consumption of a gap closing method determine how large the required computational resources must be. Therefore, we used the human dataset with real PacBio reads and compared the total CPU time and the maximum memory usage (measured as maximum resident set size across all jobs) of the gap closing tools. As shown in Table 2, DENTIST required the least amount of memory compared to all other methods, and had the second-lowest total CPU time (14 CPU days vs. 12 CPU days for LR_gapcloser). Thus, DENTIST’s speed and memory consumption is comparable to or better than other tools.
Read Coverage Analysis

Another relevant consideration is how much coverage in long reads needs to be generated to close assembly gaps and thus improve an existing fragmented assembly. A higher read coverage likely allows one to close more gaps, but also increases the sequencing costs. To evaluate how DENTIST's performance is affected by coverage, we ran our method with varying coverage of simulated PacBio reads on the D. melanogaster assembly test case. As shown in Figure 5, the number of closed gaps starts to plateau above a read coverage of about 15X. Also, above 15X the majority of gaps are closed with an accuracy ≥99%. Higher coverages increase the percent of gaps closed with ≥99% accuracy, since more reads facilitate the construction of a highly-accurate consensus sequence. In summary, while assembly improvements are also possible with low coverages of 5X or 10X, a coverage between 15X and 20X appears to be a good tradeoff between sequencing cost and power to accurately close most assembly gaps.

Application of DENTIST to a contig-only assembly

Another important feature of DENTIST is that it is able to scaffold contigs using the given long reads. In the default mode, DENTIST will only close intra-scaffold gaps that are provided in the input assembly. However, users can decide to let DENTIST additionally join different scaffolds that are bridged by reliable long read alignments, thus closing inter-scaffold gaps and increasing scaffold NG50. Additionally, DENTIST provides a “free scaffolding mode”, where it scaffolds the given contigs just using long read alignments. The latter feature is important as it extends the application range of DENTIST to contig-only assemblies, such as the more than 100 mammalian genomes produced in the Zoonomia Project that were generated mainly from short-insert library data [5].

To evaluate the scaffolding accuracy of correctly joining neighboring contigs, we discarded any scaffolding information by splitting the test assemblies into their contigs, applied DENTIST in free scaffolding mode to the four genomes using both simulated and real PacBio reads, and then analyzed the order and orientation of joined contigs. As shown in Table 3, at least 99% of DENTIST’s contig joins are in the correct order and orientation, except for Drosophila with real PacBio reads where 96.76% of the contig joins are correct. In this case, real sequencing data is used, therefore the slightly higher number of mis-assemblies (36 of 1,112) may reflect not only errors of DENTIST but also structural variations in the genome or errors in the reference assembly. Nevertheless, these results illustrate the high accuracy of the scaffolding feature of DENTIST.

Another tool that implements free scaffolding is FinisherSC, which never uses any given scaffolding information. Since FinisherSC showed a low scaffolding accuracy for Drosophila with 87% (simulated PacBio reads) and 92% (real reads) of the input contig pairs are not adjacent in the output assembly anymore (mis-assemblies that are listed as broken gaps in Table 1), we did not run further dedicated scaffolding tests.
Discussion

We have presented a novel method, DENTIST, that uses uncorrected, long sequencing reads to close gaps in fragmented assemblies. DENTIST was developed with the main goal of closing assembly gaps at a very high accuracy, for which it implements a repeat-aware read alignment step to map reads to the correct assembly loci, a consensus sequence step to obtain an accurate sequence to fill a gap, and a final validation step. Our tests using simulated and real PacBio long read data show that our method is substantially more accurate than existing tools, while achieving good sensitivity. Furthermore, DENTIST is fast and requires only moderate amounts of memory. DENTIST is also able to use long reads for closing inter-scaffold gaps and to scaffold contig-only assemblies. Together, these features make DENTIST appropriate for the task of improving the quality of hundreds of existing draft genomes with auxiliary long read data.

The accuracy of a gap closing method is mainly influenced by the accuracy of the read mappings and by the ability to determine an accurate consensus from the error-prone long sequencing reads. The latter aspect is generally challenging and even with high read coverage it is difficult to reach a desired base accuracy of Q40 (99.99% base accuracy) [1]. Therefore, de novo genome assembly from long reads includes a final “polishing” step that maps shorter Illumina reads to the finalized assembly to correct remaining base errors [1, 2]. While DENTIST already achieves a high base accuracy of the sequence inserted into gaps, this accuracy is lower than Q40. Therefore, we recommend to polish the gap-closed assembly using Illumina reads after applying DENTIST. Notably, by achieving a base accuracy of 99% or higher, DENTIST facilitates the mapping of shorter Illumina reads, which would be more difficult with the lower accuracies produced by previous methods.

To make it easy for users to run DENTIST, we used Snakemake [15] to automate the entire workflow (Figure 1) but the pipeline is built in a modularized manner and is therefore customizable. Furthermore, to enable easy application on a compute cluster without the necessity of complicated software installation steps, we provide DENTIST and all required tools in a Docker container environment that can be easily used with Snakemake’s Singularity integration [16]. The full source code is available at https://github.com/a-ludi/dentist.
Methods

Repeat masking and read mapping
Before aligning reads, DENTIST produces four types of repeat masks. First, it starts by masking low complexity regions in the given assembly using DBdust (https://dazzlerblog.wordpress.com/command-guides/dazz_db-command-guide/) in order to improve the sensitivity of the daligner alignment algorithm [17]. Second, tandem repeats are identified with datander and TANmask of the DAMASKER suite (https://dazzlerblog.wordpress.com/command-guides/damasker-commands/). Third, to identify other repetitive regions, DENTIST performs a self-alignment of the given assembly using daligner [17] and masks regions covered by at least four alignments to other genomic regions (adjustable via DENTIST’s parameter --max-coverage-self). Fourth, DENTIST creates another repeat annotation by analyzing the coverage of read alignments and marks assembly regions as repetitive that are covered by more reads than expected from the global read coverage (summed length of all long reads divided by genome size). To this end, DENTIST uses the first three repeat annotations as a soft mask and aligns all input long reads to the assembly using damapper (https://dazzlerblog.wordpress.com/2016/07/31/damapper-mapping-your-reads/), which outputs chains of local alignments arising from read artifacts such as poor quality regions or larger haplotype variations. All genomic regions covered by more than \( C_{\text{max}} \) alignments are considered repetitive. The threshold \( C_{\text{max}} \) is either given by the user via --max-coverage-reads or \( C_{\text{max}} \) is calculated from the global read coverage \( C \) (provided via --read-coverage) such that the probability of observing more than \( C_{\text{max}} \) alignments in a unique (non-repetitive) genomic region is very small (Supplementary Table 2). This probability is calculated under the assumption that the reads are sampled uniformly across the genome, implying a Poisson distribution of the number of sampled reads at any position in the genome (probability to observe \( k \) reads at any position is \( C^k \frac{e^{-C}}{k!} \)). Genomic regions with a read coverage higher than \( C_{\text{max}} \) comprise the first part of the fourth (read alignment-based) repeat annotation. To further increase the sensitivity, DENTIST searches for smaller repeat-induced local alignments. To this end, we define an alignment as proper if there are at most 100 bp (adjustable via --proper-alignment-allowance) of unaligned sequence on either end of the read. All other alignments, where only a smaller substring of the read aligns, are called improper. Improper alignments are often indicative of repetitive regions. For example, an interspersed repeat inside a long read will result in improper alignments to other genomic loci, where a similar repeat copy exists. Therefore, DENTIST considers genomic regions, where the number of improper read alignments is higher than a threshold to be repetitive. By default, this threshold equals half the global read coverage \( C \) (adjustable via --max-improper-coverage-reads). These genomic regions comprise the second part of the fourth repeat annotation.

These four repeat annotations are homogenized by transferring the annotated repeat regions to the reads using the read alignment and back again to the assembly. These homogenized, final repeat annotations are used in the next step of the pipeline.
**Alignment filtering**

To extract candidate reads that could close assembly gaps from the entire set of read alignments, it is crucial to remove potentially unreliable and irrelevant alignments. A read alignment is categorized as reliable if (1) it is proper (defined above), (2) it is strongly anchored, i.e. at least 500 bp (adjustable via \(--\text{min-anchor-length}\) of the aligned reference sequence are non-repetitive according to the homogenized repeat masks, and (3) after filtering improper alignments, every region of the aligned read must align at most to one assembly region. Finally, (4) alignments that are fully contained in a reference contig are removed because they are irrelevant for gap closing.

**Identifying closable gaps**

DENTIST identifies closable gaps by creating a so-called scaffold graph that connects input contigs based on the filtered alignments of the long reads. For every contig in the assembly, the graph has four nodes $v_{\text{begin}}^i, v_{\text{end}}^i, v_{\text{pre}}^i$, and $v_{\text{post}}^i$ representing virtual locations relative to the contig. As illustrated in Supplementary Figure 2, edges in this graph are interpreted as follows:

- $\{v_{\text{begin}}^i, v_{\text{end}}^i\}$ represents contig $i$,
- $\{v_{\text{end}}^i, v_{\text{begin}}^i\}$ for $i \neq j$ represent either reads that span contig $i$ and $j$ and thus also span the assembly gap between them, or intra-scaffold gaps (gaps between adjacent contigs) in the input assembly, and
- $\{v_{\text{pre}}^i, v_{\text{begin}}^i\}$ and $\{v_{\text{end}}^i, v_{\text{post}}^i\}$ represent reads extending the beginning or end of a contig, respectively.

Initially, the scaffold graph is populated with the contig and intra-scaffold gap edges. Then, for every read, edges of the types described above are inferred from its alignments. Note that for long reads that align to more than two consecutive contigs, we do not add the transitive edges (e.g. connecting contig $i$ to $i+2$). A list of the reads and corresponding alignments is kept for every edge such that they can be retrieved for gap closing.

**Resolving scaffolding conflicts**

The raw scaffold graph likely still contains some artifacts that need to be cleaned up. Scaffolding conflicts show up as begin nodes with more than one incoming edge or end nodes with more than one outgoing edge, meaning there is more than one way to connect the respective contig.

For assemblies with small contigs, a common conflict are small cycles resulting from reads that are long enough to completely cover one or more contigs and reach into both neighboring contigs. Depending on the local accuracy of the read, the repeat content of the intermediate contigs, and the minimum alignment length, local alignments to intermediate, small contigs may not be reliably detected, resulting in an edge that connects the neighboring contigs but
skips the intermediate one(s). However, for other reads with higher accuracy, local alignments to these small intermediate contigs can be detected, resulting in a small cycle in the scaffolding graph. To avoid mistaking these cycles for scaffolding conflicts, DENTIST searches for small cycles (up to three intermediate contigs) in the scaffold graph, identifies the “skipping” edge and aligns the reads from that edge to the intermediate (skipped) contigs with increased alignment sensitivity. If this additional step detects an alignment between the skipping read and the intermediate contig(s), the new alignments will be used to correct the graph; otherwise the read will be discarded avoiding scaffolding conflicts. This procedure effectively resolves many branching points in the scaffold graph, while preserving valuable sequence information.

The remaining scaffolding conflicts are resolved by a crude yet effective heuristic: if there are two or more spanning edges at the begin or end node of a contig, then the spanning edge with the highest number of reads will be kept, if this edge is supported by at least three times (adjustable via --best-pile-up-margin) as many reads as for the other edges; otherwise all edges in question are discarded. Additionally, to use the given scaffolding information (contig order), edges that are supported by an intra-scaffold gap in the input assembly are given a higher weight by multiplying their read number with a bonus factor (default 6, adjustable via --existing-gap-bonus).

After removing scaffolding conflicts from the graph, DENTIST discards contig-spanning edges with less than three reads, as no accurate consensus sequence for the gap can be determined from one or two reads. Also, more spanning reads give higher confidence about the correctness of the join. The minimum number of spanning reads can be increased via the parameter --min-spanning-reads to achieve a higher confidence. DENTIST also provides an “expert option” to allow gap-closing with solitary reads in which case the raw read sequence would be inserted. For valid contig-spanning edges, DENTIST puts the reads representing the spanning and extending edges into a single pile up to gather as much sequence information as possible for the consensus procedure.

**Closing the gaps**

To compute a consensus sequence for a gap, reads assigned to spanning or extending edges are cropped such that all read alignments begin and end at the same position in the reference assembly. The cropping position is chosen such that all reads still overlap the flanking contigs as much as possible to allow validation of the consensus. Cropping reduces artifacts in the subsequent pairwise alignment of the involved reads and allows easy identification of false alignments.

The cropped reads are pairwise aligned to each other using *daligner*. These read-to-read alignments often contain several local alignments, because long reads can contain regions of poor quality or larger indels and because complex gaps can result in local repeat-induced alignments. Before computing a consensus sequence, these local alignments must be filtered and “chained”. To this end, we implemented a chaining algorithm that works directly with the
alignment output produced by *daligner*. This algorithm is applied to every read-to-read alignment and reduces the alignment chaining problem to the shortest paths problem on a directed acyclic graph with node and edge weights. Every local alignment is represented by a node with a negative (beneficial) weight corresponding to the average number of base pairs covered by the local alignment of the involved reads. To determine edge weights, we define \( \text{gap}_A(x,y) \) and \( \text{gap}_B(x,y) \) as the distance between two ordered local alignments \( x \) and \( y \) on long reads \( A \) and \( B \), respectively, and \( \text{gapSizeDiff}(x,y) \) as the absolute difference between \( \text{gap}_A(x,y) \) and \( \text{gap}_B(x,y) \). Thus, \( \text{gap}_A(x,y) \) represents the number of unaligning bases in read \( A \) between \( x \) and \( y \), which is 0 if both local alignments are adjacent, and \( \text{gapSizeDiff}(x,y) \) represents the difference in the number of unaligned bases in both reads. Two nodes in the graph are connected by an edge if the respective local alignments \( x \) and \( y \) are chainable, i.e. (1) the first alignment begins strictly before the second alignment and both occur in the same orientation, (2) \( \text{gapSizeDiff}(x,y) < 1000 \) bp (adjustable via parameter \(--\text{max-indel}\) ), (3) \( \text{gap}_A(x,y) \) and \( \text{gap}_B(x,y) \) are smaller than 10,000 bp (adjustable via \(--\text{max-chain-gap}\) ), and (4) the relative overlap between the alignments, determined as the length of the overlapping region divided by the length of the smaller alignment, is \( \leq 0.3 \) (adjustable via \(--\text{max-relative-overlap}\) ). The edge then gets a positive weight, penalizing the difference between the number of unaligning bases and to smaller extent the length of the unaligning region between both local alignments as \( w_e(x,y) = \text{gapSizeDiff}(x,y) + 0.1 \cdot \max\{ |\text{gap}_A(x,y)|, |\text{gap}_B(x,y)| \} \). Maximal shortest paths in this graph constitute candidates for alignment chains. From all candidate chains, the best scoring chain(s) are selected.

After chaining, intrinsic quality values (QVs) are derived from the alignment chains using *DAScover* and *DASqv* ([https://dazzlerblog.wordpress.com/command-guides/dascrubber-command-guide/](https://dazzlerblog.wordpress.com/command-guides/dascrubber-command-guide/)). The read with the lowest number of bad QVs is chosen as the reference read, where a QV is defined as bad if it belongs to the 8% worst QVs in the pileup (adjustable via \(--\text{bad-fraction}\) ). The cleaned up alignment is used as input for *daccord* [18], which computes an accurate consensus based on the selected reference read.

Subsequently, the consensus sequence will be aligned to the flanking contigs to validate its correctness and find the exact insert sites such that the contigs are not modified when closing the gap with the consensus sequence. DENTIST allows closing gaps in three different modes. In the default mode, DENTIST closes only intra-scaffold gaps that are provided in the input assembly. In the second mode, DENTIST additionally closes gaps between different scaffolds if they are spanned by sufficient long read evidence (minimum of three spanning reads by default). In this mode, both intra- and inter-scaffold gaps are closed. In the third mode, DENTIST uses the existing scaffolding information (if available) only for conflict resolution and freely scaffolds the given contigs using the long reads. This mode enables DENTIST to also improve contig-only assemblies. Note that the selection of candidates for gap closing is only based on the evidence derived from the input read data and does not depend on the chosen mode.
Validation of closed gaps
Aiming at a high accuracy, DENTIST performs a final validation step by mapping the input reads to the gap-closed assembly. For each closed gap, DENTIST analyzes the genomic region 1000 bp (adjustable via --region-context) up- and downstream of the former gap. A closed gap is validated if there are at least three (adjustable via --min-spanning-reads) unchained read alignments spanning this region and the minimum continuous alignment coverage exceeds a user-given threshold, definable via the mandatory parameter --min-coverage-reads (alternatively, if the user provides the global long read coverage via --read-coverage and the ploidy via --ploidy, DENTIST will set the threshold to 50% of the long read coverage expected to be sequenced from a haploid locus). The continuous alignment coverage with window size \( w \) (default 500 bp; adjustable via --weak-coverage-window) at position \( x \) is defined as the number of local alignments (unchained and potentially improper) that completely cover the window \([x, x + w]\). The minimum continuous alignment coverage is then obtained by sliding the window across all positions in the genomic region defined above. For closed gaps that are not validated, DENTIST outputs the original gap (NNN…) sequence.

Evaluating the gap closing accuracy with realistic assembly gap sizes and loci
To assess the accuracy of DENTIST and compare it to other methods, we devised a realistic ground truth scenario, where the true sequence of assembly gaps is considered to be known and where assembly gaps occur at realistic genomic loci and at realistic sizes. This is important as repetitive genomic regions are the main reason for assembly gaps, thus random assembly gap placement or replacing repeats with gaps will not create a realistic setting. Each test scenario comprises (1) a high-quality reference assembly, which we consider as the ground truth, (2) a test assembly that contains assembly gaps and is input to the gap closing method, and (3) a set of long reads that were either simulated using the reference assembly or are real PacBio reads (Supplementary Table 1). Subsequently, we derived performance statistics from the output assembly of a gap closer using the same evaluation strategy.

To generate a test assembly with realistic gap sizes and loci, we aligned the reference assembly to a more fragmented, short read-based assembly of the same species and “copied” gaps from short read assembly into the respective position of the reference assembly, as depicted in Figure 2. Briefly, we aligned both assemblies using lastz [19] and constructed liftOver chains [20]. Then, we obtained the 500 bp upstream and downstream flanks of each gap in the short read assembly and used liftOver [21] (parameter -minMatch=0.8) to map these flanks to the reference assembly. Before introducing assembly gaps into the reference, we “disassembled” the reference assembly into its contigs to avoid having gaps without a known ground truth sequence. For a pair of flanks that belong to the same gap and were mapped adjacent to a reference contig, we replaced the real sequence (ground truth) between both flanks with N’s with dentist build-partial-assembly, if the gap size is at least 10 bp. We required that the reference assembly itself (which is a real assembly with gaps on its own) does not already have a gap within 3 kb of an introduced gap. It should be noted that the introduced gap size in our tests is identical to the size of the ground truth gap sequence. This gives gap
closers that take gap size into account an advantage (e.g. PBJelly, LR_gapcloser and PacBio GC) but not DENTIST, which does not use the estimated size of gaps as in reality they are often imprecise or even have a pseudo-size not related to the real gap size at all.

Since the true sequence in gaps is known, it is possible to automatically assess the performance of the gap closing tools. To this end, we first identify the original contigs of the test assembly in a gap-closed assembly by searching for exact and unique matches. Duplicated contigs that already have more than one exact match within the test assembly are excluded. We noticed that some gap closing tools do not only fill in gaps but also modify the input contigs. We handled this unexpected behavior in two ways. First, before searching for exact matches, we remove a given number of base pairs from the flanks of the contig of the test assembly, thus allowing the gap closer to modify contig flanks. This number of base pairs in controlled by --crop-ambiguous (default 100bp) when searching for duplicate contigs and by --crop-alignment when searching for contigs in the gap-closed assembly. Second, after identifying exact matches, we optionally conduct a second search for the remaining contigs, allowing for up to 1.5% mismatches but requiring that a contig fully and uniquely aligns to some part of the result assembly. Consequently, original contigs that are substantially modified by a gap closing method may not be detected, which highlights an unwanted behavior of the respective method.

After determining the locations of the test contigs, each gap is evaluated according to the following rules. A gap is considered closed if the locations of both contigs surrounding the gap in the test assembly are mapped to a single contig in the gap-closed assembly. In this case, the sequence identity between the known gap sequence and the inserted gap sequence is calculated. A gap is considered unclosed if the locations of both contigs surrounding the gap in the test assembly are known and the contigs are mapped to different but adjacent contigs in the gap-closed assembly, i.e. a gap remains. A gap is considered unknown if none of the flanking contigs are found or at least one flanking contig cannot be located uniquely. These gaps are excluded from further analysis, as we cannot accurately determine the inserted gap sequence. If neither of the above cases is true, the gap is said to be broken. This is the case if both flanking contigs could be located but are not adjacent anymore, i.e. the contigs are misassembled. This procedure allows to obtain an accurate assessment of the performance of each gap closing tool in an automated fashion.

Long reads used in our tests were either real PacBio reads (Supplementary Table 1) or reads that were sampled (simulated) from the reference assembly adding a typical PacBio base and indel error profile using “simulator -m25000 -s125000 -e.13 -rSEED” (https://dazzlerblog.wordpress.com/command-guides/dazz_db-command-guide/). The seed used for simulating reads is listed in Supplementary Table 1. For real PacBio reads, we provided all subreads from all wells to the gap closing tools.
Running gap closing methods
All tools were called with the default or recommended parameters, except for parameters that do not influence the output such as the number of parallel threads. For finisherSC.py, we used parameters “--fast True --large True” to make the Drosophila application feasible in terms of runtime and memory requirements.

The results were evaluated using “dentist check-results” with parameter “--crop-ambiguous=300” to ignore 300 bp from the contig flanks when searching for duplicate contigs. To evaluate PBJelly, parameter “--crop-alignment=100” was used in addition to ignore 100 bp from the contig flanks in the contig identification step. To evaluate LR_gapcloser, which can modify not only the original contig flanks, we additionally specified parameters “--crop-alignment=300” and “--recover-imperfect-contigs” to enable contig identification with up to 1.5% mismatches. The increase in contiguity of the gap-closed assembly was measured by the percentage of closed gaps and the increase in contig NG50. The correctness of closed gap sequences was measured as the sequence identity between the (known) ground truth and the inserted sequence for each gap. These values are presented in three forms: as a distribution binned in six intervals: [0, 0.7), [0.7, 0.9), [0.9, 0.95), [0.95, 0.99), [0.99, 1.0), and {1.0}, as the arithmetic mean over all the sequence identities, and as the weighted arithmetic mean using the true gap sizes as weights.

For tests with simulated reads, we also used “dentist find-closable-gaps” to determine which gaps are closable, as the true origin of each read is known. We defined a gap as closable if and only if at least three reads span the gap and 500 bp on either side.

Evaluating DENTIST’s scaffolding mode
To evaluate the scaffolding accuracy of correctly joining neighboring contigs, we discarded any scaffolding information from the test assemblies by splitting them into contigs, and applied DENTIST in free scaffolding mode (parameter --join-policy=contigs). We then analyzed the scaffolding information provided by DENTIST (parameter --scaffolding) for each join separately. We counted a join as correct if the involved contig numbers are ascending (descending for reverse-complement orientation) and the contigs are in the correct orientation. In case a join skipped some contigs (e.g. contigs i and i + 1 + x with x ≥ 1 were joined), we counted it still as correct if we could successfully map the skipped contig(s) in the correct order and orientation to the inserted sequence representing the larger closed gap. Since DENTIST might also join the end-contigs of input scaffolds where we do not know the true scaffolding, we excluded these “novel” joins from the analysis. All other joins are counted as incorrect.
Data and code availability
The DENTIST code including a Docker container and Snakemake workflow is available at https://github.com/a-ludi/dentist. All data, including the reference and test assemblies with introduced gaps and their true sequence as valuable data for future method comparisons, is available at https://bds.mpi-cbg.de/hillerlab/DENTIST/.

Competing interests
The authors have no competing interests.

Acknowledgment
We thank the Computer Service Facilities of the MPI-CBG for their support. This work was supported by the Max Planck Society, the Federal Ministry of Education and Research (grant 01IS18026C), and the LOEWE-Centre for Translational Biodiversity Genomics (TBG) funded by the Hessen State Ministry of Higher Education, Research and the Arts (HMWK).
References

1. Rhie A, McCarthy SA, Fedrigo O, Damas J, Formenti G, Koren S, Uliano-Silva M, Chow W, Fungtammasan A, Gedman GL, et al: Towards complete and error-free genome assemblies of all vertebrate species. bioRxiv 2020;2020.2005.2022.110833.

2. Jebb D, Huang Z, Pippel M, Hughes GM, Lavrichenko K, Devanna P, Winkler S, Jermiin LS, Skirmuntt EC, Katzourakis A, et al: Six reference-quality genomes reveal evolution of bat adaptations. Nature 2020, 583:578-584.

3. Warren WC, Harris RA, Haukness M, Fiddes IT, Murali SC, Fernandes J, Dishuck PC, Storer JM, Raveendran M, Hillier LW, et al: Sequence diversity analyses of an improved rhesus macaque genome enhance its biomedical utility. Science 2020, 370.

4. Rice ES, Green RE: New Approaches for Genome Assembly and Scaffolding. Annu Rev Anim Biosci 2019, 7:17-40.

5. Zoonomia C: A comparative genomics multitool for scientific discovery and conservation. Nature 2020, 587:240-245.

6. Feng S, Stiller J, Deng Y, Armstrong J, Fang Q, Reeve AH, Xie D, Chen G, Guo C, Faircloth BC, et al: Dense sampling of bird diversity increases power of comparative genomics. Nature 2020, 587:252-257.

7. Miga KH, Koren S, Rhie A, Vollger MR, Gershman A, Bzikadze A, Brooks S, Howe E, Porubsky D, Logsdon GA, et al: Telomere-to-telomere assembly of a complete human X chromosome. Nature 2020, 585:79-84.

8. English AC, Richards S, Han Y, Wang M, Vee V, Qu J, Qin X, Muzny DM, Reid JG, Worley KC, Gibbs RA: Mind the gap: upgrading genomes with Pacific Biosciences RS long-read sequencing technology. PLoS One 2012, 7:e47768.

9. Lam KK, LaButti K, Khalak A, Tse D: FinisherSC: a repeat-aware tool for upgrading de novo assembly using long reads. Bioinformatics 2015, 31:3207-3209.

10. Xu GC, Xu TJ, Zhu R, Zhang Y, Li SQ, Wang HW, Li JT: LR_Gapcloser: a tiling path-based gap closer that uses long reads to complete genome assembly. Gigascience 2019, 8.

11. Kosugi S, Hirakawa H, Tabata S: GMcloser: closing gaps in assemblies accurately with a likelihood-based selection of contig or long-read alignments. Bioinformatics 2015, 31:3733-3741.

12. Xu M, Guo L, Gu S, Wang O, Zhang R, Peters BA, Fan G, Liu X, Xu X, Deng L, Zhang Y: TGS-GapCloser: A fast and accurate gap closer for large genomes with low coverage of error-prone long reads. Gigascience 2020, 9.

13. Lu P, Jin J, Li Z, Xu Y, Hu D, Liu J, Cao P: PGcloser: Fast Parallel Gap-Closing Tool Using Long-Reads or Contigs to Fill Gaps in Genomes. Evol Bioinform Online 2020, 16:1176934320913859.

14. Wenger AM, Peluso P, Rowell WJ, Chang PC, Hall RJ, Concepcion GT, Ebler J, Fungtammasan A, Kolesnikov A, Olson ND, et al: Accurate circular consensus long-read sequencing improves variant detection and assembly of a human genome. Nat Biotechnol 2019, 37:1155-1162.

15. Koster J, Rahmann S: Snakemake—a scalable bioinformatics workflow engine. Bioinformatics 2012, 28:2520-2522.

16. Kurtzer GM, Sochat V, Bauer MW: Singularity: Scientific containers for mobility of compute. PLoS One 2017, 12:e0177459.

17. Myers G: Efficient Local Alignment Discovery amongst Noisy Long Reads. In; Berlin, Heidelberg. Springer Berlin Heidelberg; 2014: 52-67.

18. Tischler G, Myers EW: Non Hybrid Long Read Consensus Using Local De Bruijn Graph Assembly. bioRxiv 2017:106252.

19. Harris RS: Improved pairwise alignment of genomic DNA. The Pennsylvania State University, 2007.

20. Kent WJ, Baertsch R, Hinrichs A, Miller W, Haussler D: Evolution's cauldron: duplication, deletion, and rearrangement in the mouse and human genomes. Proc Natl Acad Sci U S A 2003, 100:11484-11489.

21. Hinrichs AS, Karolchik D, Baertsch R, Barber GP, Beijerano G, Clawson H, Diekhans M, Furey TS, Harte RA, Hsu F, et al: The UCSC Genome Browser Database: update 2006. Nucleic Acids Res 2006, 34:D590-598.
**Figures**

**Figure 1:** Overview of the assembly gap closing pipeline implemented in DENTIST.

**Figure 2:** Realistic ground truth scenarios are created by “copying” gaps from a short read assembly to a high-quality reference assembly. Horizontal lines represent contigs, blue boxes represent genomic regions of the reference assembly, which were replaced with assembly gaps (N’s). The white box in the reference assembly represents an assembly gap in the reference, which is not copied to the test assembly.
Figure 3: Comparison of accuracy and sensitivity of gap closing methods.

(A, B) Bar charts show the percent of gaps that are closed by different methods using simulated (A) or real (B) PacBio reads and a breakdown of how identical the inserted sequence is to the ground truth sequence.

(C, D) NG(x) plots show which percent of the assembly consists of contigs of a certain minimum size (Y-axis). The contig NG50 and NG90 values are indicated by vertical dashed lines. The test assembly, which contains introduced assembly gaps, is shown as a dashed
grey line, the ground truth assembly, representing the maximally-possible contiguity if all introduced gaps are closed, is shown as a solid grey line. Due to excessive memory consumption, we could run FinisherSC only on the *Drosophila* genome and this method produced many broken gaps (likely representing mis-assemblies, as the input contigs are not adjacent anymore in the output assembly). PacBio GC requires real read data as input.

Figure 4: Comparison of base identity and gap size using the human assembly and real PacBio reads. (A) Gaps, which are closed at an accuracy of <98% by DENTIST, tend to be short. Many of these gaps contain simple repeats such as homopolymer runs, for which long reads have a higher error rate. In contrast, gaps closed at a lower accuracy by PBJelly (B), LR_gapcloser (C) and PacBio GC (D) are more evenly distributed in size. Furthermore, these three methods rarely close larger gaps (>1000 bp) with near 100% accuracy. For PBJelly and LR_gapcloser, there is a trend that the longer the gaps, the lower is the accuracy.
Figure 5: Coverage analysis by applying DENTIST to *Drosophila* with varying coverage of simulated PacBio reads.

The figure compares contig NG50 (top), average identity of the closed gaps (middle) and the number of closed gaps together with a breakdown of their sequence identity (bottom). The number of closed gaps starts to plateau above a read coverage of about 15X (indicated by a vertical dashed line).
For DENTIST, we did not count the runtime of local Snakemake rules which consume very adjacent anymore after g average weighted by the true gap length. Gaps are broken if the flanking contigs are not real

Table 1: Comprehensive benchmarking of gap closing methods.

We define %ID as the average across all sequence identities of closed gaps and w%ID as the average weighted by the true gap length. Gaps are broken if the flanking contigs are not adjacent anymore after gap closing, indicating a mis-assembly. For simulated reads, we label a gap as closable if there are at least three simulated reads that span the gap and at least 500 bp of its flanks. The contig NG50 value of the test (input) and output assembly is given on the right; maximum contig NG50 refers the value if all introduced gaps would be closed. Best values in each category are in bold.

Table 2: Comparison of runtime and maximum memory consumption on the human assembly with real PacBio reads.

For DENTIST, we did not count the runtime of local Snakemake rules which consume very little resources. For PacBio GC, we excluded resources needed for the expensive pre- and post-processing of the assembly, but included the essential calls of samtools faidx, pbindex, pbalign and variantCaller. GiB = Gibibyte (1 073 741 824 ≈ 230 bytes).
### Table 3: Accuracy of DENTIST in scaffolding contigs with long reads.

In these tests, we applied DENTIST in free scaffolding mode, where it infers the contig order and orientation solely from alignments of long reads, simulating the application to a contig-only assembly. For Drosophila and real reads, DENTIST made 8 additional joins between pairs of contigs located on scaffold ends, thus closing an inter-scaffold gap. As it is not known whether these joins are correct, they were not included in the breakdown of total joins.

| Species   | Read type | contig joins | % correct joins |
|-----------|-----------|--------------|-----------------|
|           |           | total  | correct | incorrect |             |
| Drosophila| simulated| 1,278  | 1,268   | 10        | 99.22%     |
|           | real      | 1,112  | 1,076   | 36        | 96.76%     |
| Arabidopsis| simulated| 233    | 231     | 2         | 99.14%     |
|           | real      | 164    | 163     | 1         | 99.39%     |
| Hummingbird| simulated| 36,868 | 36,664  | 204       | 99.45%     |
|           | real      | 34,489 | 34,299  | 190       | 99.45%     |
| Human     | simulated| 4,612  | 4,566   | 46        | 99.00%     |
|           | real      | 4,000  | 3,988   | 12        | 99.70%     |