Assemblathon 1: A competitive assessment of de novo short read assembly methods


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1. Abstract

Low cost short read sequencing technology has revolutionised genomics, though it is only just becoming practical for the high quality de novo assembly of a novel large genome. We describe the Assemblathon 1 competition, which aimed to comprehensively assess the state of the art in de novo assembly methods when applied to current sequencing technologies. In a collaborative effort teams were asked to assemble a simulated Illumina HiSeq dataset of an unknown, simulated diploid genome. A total of 41 assemblies from 17 different groups were received. Novel haplotype aware assessments of coverage, contiguity, structure, base calling and copy number were made. We establish that within this benchmark (1) it is possible to assemble the genome to a high level of coverage and accuracy, and that (2) large differences exist between the assemblies, suggesting room for further improvements in current methods. The simulated benchmark, including the correct answer, the assemblies and the code that was used to evaluate the assemblies is now public and freely available from http://www.assemblathon.org/.

2. Introduction
Sequence assembly is the problem of merging and ordering shorter fragments, termed “reads,” sampled from a set of larger sequences in order to reconstruct the larger sequences. The output of an assembly is typically a set of “contigs,” which are contiguous sequence fragments, ordered and oriented into “scaffold” sequences, with gaps between contigs within scaffolds representing regions of uncertainty.

There are numerous subclasses of assembly problem that can be distinguished by, amongst other things, the nature of (1) the reads, (2) the types of sequences being assembled and (3) the availability of homologous (related) and previously assembled sequences, such as a reference genome or the genome of a closely related species (Pop & Salzberg, 2008) (Trapnell & Salzberg, 2009) (Chaisson, Brinza, & Pevzner, Genome Res, 2009). In this work we focus on the evaluation of methods for de novo genome assembly using low cost “short read” technology, where the reads are comparatively short in length but large in number, the sequences being assembled represent a novel diploid genome and the nearest homologous genome to that being assembled is significantly diverged.

In bioinformatics, the reads used in an assembly are derived from an underlying sequencing technology. For a recent review of sequencing technologies see Metzker (2010). For the assembly problem there are a number of key considerations, notably (1) the length of the reads, (2) the error characteristics of the reads, (3) if and how the reads are “paired”, i.e. where reads are produced in pairs separated by an approximately fixed length spacer sequence, and finally (4) the number of reads produced for a given cost.

Sanger sequencing (Sanger, Nicklen, & Coulson, 1977) produces relatively long reads, typically between 300 to 1000 base pairs (bp) in length, with a low error rate, but which are comparatively expensive to produce. After relying primarily on Sanger sequencing for decades the field of sequencing has recently witnessed a diversification of competing technologies ((Margulies, et al., 2005), (Bentley, 2006), (Pandey, Nutter, & Prediger, 2008), (Eid, et al., 2009), (Pourmand, et al., 2006)) and a rapid rate of overall change. One direction of this development has been a move towards shorter reads, often less than or equal to 150 bp, but at a much lower cost for a given volume of reads ((Bentley, 2006), (Pandey, Nutter, & Prediger, 2008), (Pourmand, et al., 2006)).

As the field of sequencing has changed so has the field of sequence assembly, for a recent review see Miller, Koren & Sutton (2010). In brief, using Sanger sequencing, contigs were initially built using overlap or string graphs (Myers E., 2005) (or data structures closely related to them), in tools such as Phrap (http://www.phrap.org/), GigAssembler (Kent & Haussler, 2001), Celera
(Myers, et al., 2000) (Venter, et al., 2001), ARACHNE (Batzoglou, et al., 2002), and Phusion (Mullikin & Ning, 2003), which were used for numerous high quality assemblies such as human (Lander, et al., 2001) and mouse (Mouse Genome Sequencing Consortium, et al., 2002). However, these programs were not generally efficient enough to handle the volume of sequences produced by the next generation sequencing technologies, spurring the development of a new generation of assembly software.

While some maintained the overlap graph approach, e.g. Edena (Hernandez, François, Farinelli, Osterås, & Schrenzel, 2008) and Newbler (http://www.454.com/), others used word look-up tables to greedily extend reads, e.g. SSAKE (Warren, Sutton, Jones, & Holt, 2007), SHARCGS (Dohm, Lottaz, Borodina, & Himmelbauer, 2007), VCAKE (Jeck, et al., 2007) and OligoZip (http://linux1.softberry.com/berry.phtml?topic=OligoZip). These word look-up tables were then extended into de Bruijn graphs to allow for global analyses (Pevzner, Tang, & Waterman, 2001), e.g. Euler (Chaisson & Pevzner, Genome Res, 2008), AllPaths (Butler, et al., 2008) and Velvet (Zerbino & Birney, 2008). As projects grew in scale further engineering was required to fit large whole genome datasets into memory ((ABySS (Simpson, Wong, Jackman, Schein, Jones, & Birol, 2009), Meraculous (in submission)), (SOAPdenovo (Li, et al., 2010), Cortex (in submission)). Now, as improvements in sequencer technology are extending the length of “short reads”, the overlap graph approach is being revisited, albeit with optimized programming techniques, e.g. SGA (Simpson & Durbin, Bioinformatics, 2010), as are greedy contig extension algorithms, e.g. PRICE (http://derisilab.ucsf.edu/software/price/index.html), Monument (http://www.irsia.fr/symbiose/people/rchkhi/monument.html).

In general, most sequence assembly programs are multi stage pipelines, dealing with correcting measurement errors within the reads, constructing contigs, resolving repeats (i.e. disambiguating false positive alignments between reads) and scaffolding contigs in separate phases. Since a number of solutions are available for each task, several projects have been initiated to explore the parameter space of the assembly problem, in particular in the context of short read sequencing ((Phillippy, Schatz, & Pop, 2008), (Hubisz, Lin, Kellis, & Siepel, 2011), (Alkan, Sajjadian, & Eichler, 2011), (Narzisi & Mishra, 2011), (Zhang, Chen, Yang, Tang, Shang, & Shen, 2011) and (Lin, Li, Shen, Zhang, Papasian, & Deng, 2011)). In this work we are concerned with evaluating assembly programs as whole, with the aim of comprehensively evaluating different aspects of assemblies.

It is generally the case that the right answer to an assembly problem is unknown. Understandably therefore, a common method for assessing assembly quality has been the calculation of length summary statistics on the produced scaffold and contig sequences. Such metrics include various
weighted median statistics, such as the N50 defined below, as well as the total sequence lengths and total numbers of sequences produced ((Lindblad-Toh, et al., 2005), (Ming, et al., 2008), (Liu, et al., 2009), (Church, et al., 2009), (Li, et al., 2010), (Locke, et al., 2011) and (Colbourne, et al., 2011)).

Other methods have been proposed for evaluating the internal consistency of an assembly, for example, by analysing the consistency of paired reads, as in the clone-middle plot (Huson, Halpern, Lai, Myers, Reinert, & Sutton, 2001), by looking for variations in the depths of read coverage supporting a constructed assembly (Phillippy, Schatz, & Pop, 2008) and looking at haplotype inconsistency (Lindblad-Toh, et al., 2005).

To assess accuracy, assemblies may be compared to finished sequences derived from independent sequencing experiments or to sequences held out of the assembly process. For the dog genome, nine bacterial artificial chromosomes (BACs) were sequenced to finishing standards and held out of the assembly (Lindblad-Toh et al. 2005), for the panda genome, which was primarily an Illumina assembly, extra Sanger sequencing of BACs was performed (Li et al. 2010). Additionally, if genetic mapping data is available such information can also be used to assess scaffold quality, e.g. Church et al. (2009), which used a combination of linkage, radiation hybrid and optical maps. Church et al. also demonstrate that transcriptome (the set of RNA molecules for a given cell type) information, if available, can also be used to assess the validity of a genomic assembly by checking the extent to which the assembly recapitulates the transcriptome.

When a reference genome or sequence is available a comparison between the assembly and reference can be performed. This has previously been accomplished by studies using several different genome alignment methods, including BLAST (e.g. Zang et al. 2011), LASTZ (e.g. (Hubisz, Lin, Kellis, & Siepel, 2011)) and Exonerate (Zerbino & Birney 2008, Hernandez et al. 2008). Given such an alignment, most simply, the proportion of a reference's coverage can be reported (Li, et al., 2010) (Zhang, Chen, Yang, Tang, Shang, & Shen, 2011). Notably, Gnerre et al. (2011) compared novel short-read assemblies to the human and mouse reference genomes, and performed a comprehensive set of analyses that encompassed coverage, contig accuracy and the long range contiguity of scaffolds. Related to the work described here, Butler et al. (2008) described a graph based pattern analysis using an assembly to reference alignment.

Comparison can also be made to a well-sequenced, related species. This can be done using the complete genomic sequence of an out group, for example, Meader et al. (Meader, Hillier, Locke, Ponting, & Lunter, 2010) presented an assessment method based on patterns of insertions and
deletions (indels) in closely related inter-species genome alignments. Alternatively, specific genomic features can be studied, for example Parra et al. examined the fraction of “core genes,” those present in all genomes, that could be identified in draft genome assemblies (Parra, Bradnam, Ning, Keane, & Korf, 2009).

Simulation has been a mainstay of genome assembly evaluation since assembly methodology was first developed and with few exceptions (e.g. (Maccallum, et al., 2009), (Gnerre, et al., 2011)) is de rigueur when introducing new de novo assembly software (e.g. (Myers, et al., 2000), (Lander, et al., 2001), (Venter, et al., 2001), (Batzoglou, et al., 2002), (Warren, Sutton, Jones, & Holt, 2007), (Jeck, et al., 2007), (Dohm, Lottaz, Borodina, & Himmelbauer, 2007), (Chaisson & Pevzner, Genome Res, 2008), (Zerbino & Birney, 2008), (Butler, et al., 2008) etc). In this work we have also chosen to use simulations, utilising the new Evolver genome evolution simulation tool (http://www.drive5.com/evolver/) to produce a simulated diploid genome with parameters that approximate that of a vertebrate genome, though at ~1/10th the scale.

From this novel genome we simulate reads, modelling an Illumina sequencing run, using a newly developed read simulator. Assembly teams were asked to assemble this novel genome blind and we present an analysis of the resulting assemblies. By using simulation we know a priori the haplotype relationships; by a process of multiple sequence alignment (MSA) we assess the relationships between the assemblies and the original haplotypes of our simulation. This novel process allows us evaluate haplotype specific contributions to the assemblies. Additionally, as a positive control for our results, we assess the generated assemblies using more traditional BLAST (Zang et al. 2011), (Hubisz, Lin, Kellis, & Siepel, 2011)) methods, and support our assessments by making all the code and data from our assessments public and freely available (http://compbio.soe.ucsc.edu/assemblathon1/, http://www.assemblathon.org/).

3. Results

We start by giving an overview of the Assemblathon 1 dataset and its generation. We then describe the assemblies before giving the results of different evaluations.

3.1 Genome simulation

Rather than use an existing reference genome for assessment, we opted to simulate a novel genome. We did this primarily for three reasons. Firstly, it gave us a genome that had no reasonable homology to anything other than out-group genomes that we generated and provided
to assemblers. This allowed for a fair, blind test in which none of the assembly contributors had access to the underlying genomes during the competition. Secondly, we were able to precisely tailor the proportions of the simulated genome to those desired for this experimental analysis, i.e. to limit the size of the genome to less than that of a full mammalian genome and thus allow the maximum number of participants, while still maintaining a size that posed a reasonable challenge.

Thirdly, we could simulate a diploid genome; we know of no existing diploid dataset (simulated or real) in which the contributions of the two haplotypes are precisely and fully known. This allowed us to assess a heretofore-unexplored dimension of assembly assessment.

To simulate the genome we used the Evolver suite of genome evolution tools. Evolver simulates the forward evolution of multi-chromosome haploid genomes and includes models for evolutionary constraint, protein codons, genes and mobile elements.

The input genome for the simulation, termed the root genome, was constructed by downloading the DNA sequence and annotations (see methods) for human chromosome 13 (hg18/NCBI36, 95.6 non-N megabases (Mb)) from the UCSC Table Browser (Fujita, et al., 2011) and dividing it into four chromosomes of approximately equal length. Figure 1 shows the phylogeny used to generate the simulated genomes, with branch lengths to scale. We first evolved the root genome for ~200 million years (my) to generate the most recent common ancestor (MRCA) of the final leaf genomes. We performed this long burn-in on the genome in order to reshuffle the sequence and annotations present, thereby preventing simple discovery of the source of the root genome. The simulation then proceeded along two independent lineages, generating both $\alpha$ and $\beta$, each ~50 my diverged from the MRCA. Finally, in both lineages we split the evolved genome into two sub-lineages, termed haplotypes, and evolved these sub-lineages for a further ~1 my, to produce a pair of diploid genomes $\alpha_{1,2}$ and $\beta_{1,2}$, each with a degree of polymorphism. The $\alpha_{1,2}$ genome’s haplotypes, $\alpha_1$ and $\alpha_2$, each had three chromosomes and both haplotypes were 112.5 Mb in total length with chromosome lengths of 76.3, 18.5 and 17.7 Mb.

The diploid $\alpha_{1,2}$ genome was used as the target genome for the assembly. The $\beta_{1,2}$ genome’s haplotypes, their common ancestor, $\beta$ and their annotations were provided to the assemblers as an out-group. Relatively few assemblers (see Table 1) reported using these sequences to assist in the assembly process.

Table 2A provides a count of some of the events that took place along particular branches in the phylogenetic tree during the course of the simulation. Table 2B provides a summary of the
pairwise differences between the $\alpha_1$ and $\alpha_2$ haplotypes and Supplementary Figure 1 shows a dot-plot of their alignment. Supplementary Figures 2 and 3 show the length distribution of annotations for the root, MRCA, internal node and leaf genomes, demonstrating these annotations remained approximately static over the course of the simulation. We examined repeat content of the simulated genomes (see Table 2 legend) and found a comparable portion of annotated repeats to that in the original human chromosome 13, but a reduction of slightly more than half in the proportion of repetitive 100-mers (Supplementary Figure 4). The simple substitution model used by Evolver, which fails to capture some of the higher order dependencies in substitution patterns that made the original human DNA sequence more repetitive, likely explains this latter observation.

3.2 Read Simulation

As mentioned, there are many competing technologies now available for sequencing, giving us many possible options in designing the datasets for the first Assemblathon. However, we opted to simulate just one combined short read dataset, with multiple read libraries, for the Illumina Hi-seq platform (Illumina, Illumina HiSeq 2000), which is the current market leader for low cost de novo sequencing on this scale. The advantage of this was chiefly (1) the avoidance of fragmentation in the entries to the Assemblathon, thereby preventing categories with few or just one entry, and (2) the ability to assess all the submitted assemblies with common sets of evaluations.

We needed a program that would generate short reads and model sources of error that the Illumina protocols introduce. As we knew of no existing software that was capable of this (see methods for a discussion of existing read simulators), we wrote our own short read simulator called SimSeq (https://github.com/jstjohn/SimSeq).

Abstractly, reads were sampled from the genome using one of two types of Illumina paired read strategy, so called ‘paired-end’ and ‘mate-pair’ strategies, after which an error profile was applied to each read in its proper orientation. In addition to generating reads from the target $\alpha_{1,2}$ genome, three copies of an Escherichia coli sequence (gi 312944605) were added to the two haplotype sequences to yield a ~5% bacterial contamination rate. Bacterial sequence was included as an attempt model the sort of contamination occasionally present in data from sequencing centres, though the specific choice of E.coli and the 5% level were arbitrary. Participants in the contest were notified that some bacterial contamination was present in the data, though they were not told about its precise nature nor explicitly told to remove it.
Multiple libraries were generated for both the paired-end and mate-pair strategies. Paired-end libraries with 200 and 300 bp inserts contributed 80x, mate-pair libraries with separations of 3 kilobases (kb) and 10 kb contributed a further 40x, giving a total coverage of 120x for the sample. Removing contamination reads gave an overall coverage of ~55x per haplotype.

A detailed description of the simulation method, the types of errors simulated, and the simulator's limitations are given in the methods. Importantly, due to human error, the error model was mistakenly reversed along the reads. This resulted in bases with a slightly higher error rate tending to appear towards the beginning of the reads rather than towards the end of reads (see Supplementary Figure 5). This issue only manifests itself if the reads are treated asymmetrically; we surveyed participants on this matter and only one group, L'IRISA, indicated that their methodology was possibly harmed more than other methods due to the mistake.

### 3.3 Assemblies

The competition started in January 2011 and teams were given just over a month to submit their assemblies. Teams were allowed to submit up to five separate assemblies for consideration. Additionally, assemblies were created by the organisers with popular assembly programs, using default parameters, as a way of comparing naively generated assemblies with those that were contributed by independent groups. Table 1 lists the evaluated assemblies, the main program used to generate them and the groups that contributed them (see Supplementary section 8.2 for detailed information on submissions). In total there were 59 assemblies, with 41 independently contributed by 17 different groups using 15 different assembly programs and 18 generated by the organisers using three popular programs.

### 3.4 Evaluations

We assessed all the contributed assemblies, full results for which can be found in the supplementary material. However, to make the presentation succinct we choose to present only the “top” assembly from each group in the following evaluations. To enable this we created a ranking of the assemblies (see Table 3, Supplementary Table 1), using the evaluations described below, and selected the assembly from each group with the top overall ranking for inclusion. Full results for each evaluation on every assembly in the main text can be found in the supplementary material.
3.4.1 N50 and NG50

A commonly used metric to assess assemblies is the N50 statistic. The N50 of an assembly is a weighted median of the lengths of the sequences it contains, equal to the length of the longest sequence $s$ such that the sum of the lengths of sequences greater than or equal in length to $s$ is greater than or equal to half the length of the genome being assembled. As the length of the genome being assembled is generally unknown, the normal approximation is to use the total length of all of the sequences in an assembly as a proxy for the denominator. We follow this convention for calculating N50, but additionally we define the $NG50$ (G for genome). The NG50 is identical to N50, except that we estimate the length of the genome being assembled as being equal to the average of the length of the two haplotypes, $\alpha_1$ and $\alpha_2$. Contig N50s and NG50s, where the sequences are the set of assembly contigs, and scaffold N50s and NG50s, where the sequences are the set of assembly scaffolds, are shown in Figure 2, Supplementary Figure 6 and Supplementary Table 2.

The total span of most of the submitted assemblies was slightly larger than the haploid genome size, primarily because of the degree of polymorphism of the two haplotypes. Thus the assembly-specific N50s are in general smaller than the NG50s, with the median absolute difference between contig NG50 and contig N50 being 599 bp (7.7%), and the median absolute difference between scaffold NG50 and scaffold N50 being 1942 bp (3.6%). These differences are quite small, though not negligible in every case, for example the CSHL assembly has a scaffold NG50 ~800 kb (31.6%) longer than scaffold N50.

3.4.2 Multiple Sequence Alignment

While N50 statistics give a sense of the scale and potential contiguity of an assembly they say nothing necessarily about the underlying coverage or accuracy of an assembly. To compare each assembly with the simulated genome and bacterial contamination we constructed a multiple sequence alignment (MSA). The sequence inputs to the MSA were the two haplotypes, the bacterial contamination and the scaffolds of the assembly. To generate the MSA we used an adapted version (see methods) of the newly developed Cactus alignment program (Paten, Earl, Nguyen, Diekhans, Zerbino, & Haussler, 2011), a new MSA program able to handle rearrangements, copy number changes (duplications) and missing data. The result of this alignment process was, for each assembly, a high specificity map of the alignment of the assembly to the two haplotypes and the bacterial contamination.
As we aligned both the bacterial contamination and the two haplotypes together we used the hypothetical existence of any alignments between the haplotypes and the bacterial contamination as a negative control for non-specific alignment. We did not observe any such alignments. As an additional control we replicated a similar, confirmatory analysis using a simple BLAST (Altschul, Gish, Miller, Myers, & Lipman, 1990) strategy, details of which can be found in Supplementary section 7.1 and references to which are made below.

### 3.4.3 Coverage

An MSA can be divided up into columns, each of which represents a set of individual base pair positions in the input sequences that are considered homologous. We call columns that contain positions within the haplotypes haplotype columns. We define the overall coverage of an MSA as the proportion of haplotype columns that contain positions from the assembly. Similarly we can define the coverage of X, where X is a specific haplotype or the bacterial contamination, as the proportion of columns containing positions in X that also contain positions from the assembly.

Table 4 shows the overall, haplotype specific, and bacterial contamination specific coverage. There is very little difference between the specific haplotype’s coverage and the overall coverage, and indeed little difference between many of the assemblies. The highest overall coverage was the BGI assembly with 98.8%, but nearly all assemblies performed well in this metric with even the assembly with 14th highest coverage, the CRACS assembly, providing 96.3% coverage. However, there were huge differences in the coverage of the bacterial contamination (Supplementary Figures 7 and 8), with many groups opting successfully to completely filter it out. For example, the BGI assembly had no coverage of the bacterial sequence, while the ASTR assembly had 100.0% coverage of the bacterial sequence.

### 3.4.4 Blocks and Contig Paths

Within an MSA a block is a maximal gapless alignment of a set of sequences, and is therefore composed of a series of contiguous columns. The length of a block is equal to the number of columns that it contains. We can use the block structure to define the block NG50, which is exactly like the NG50, except that we use the distribution of block lengths rather than sequence lengths. Supplementary Figures 9 and 10 show block coverage across the haplotypes. Alignment of sequences that are very closely related are likely to contain fewer blocks with a greater base pair length than sequences that are significantly diverged from one another. Unfortunately, the two simulated haplotypes are sufficiently polymorphic with respect to one another that the block
NG50 of an alignment of just the two haplotypes is ~4 kb. As this length is much less than the
length of many sequences in the assemblies, assessing an assembly requires methods that do
not penalise the reconstruction of haplotype specific polymorphisms. This is evident by looking at
Figure 2, which shows that block NG50 is poorly discriminative. See Supplementary section 7.1.1
and Supplementary Figure 11 for supporting BLAST based analysis.

To extend our analysis we use a graph theoretic model of the alignments, which we now describe
in overview. An MSA can be described as a graph, and we call the simplest such graph an
adjacency graph. A formal description of the adjacency graph used here can be found in (Paten,
et al., 2011), it is closely related to the similarly name graph introduced in (Bergeron, Mixtacki, &
Stoye, 2006), but also to a directed bigraph representation of a de Bruijn graph used in assembly
(Medvedev & Brudno, 2009) and the multiple breakpoint graph used in the study of genome
rearrangements (Alekseyev & Pevzner, 2009).

An adjacency graph $G$ contains two kinds of edges, block edges, which represent the gapless
blocks of the alignment, and adjacency edges, which represent collections of connections
between the ends of segments of DNA. The nodes in the graph represent the ends of blocks of
aligned sequences. Figure 3 illustrates an example.

Each edge in $G$ is labelled with the sub-sequences it represents, called segments, thus it is
possible to discern if the edge represents segments in the haplotypes, the assembly, the bacterial
contamination or some combination. As previously stated, no edges are contained in $G$ that
represent segments in both the haplotypes and the bacterial contamination.

Within $G$ a sequence is represented as a path of alternating adjacency and block edges, termed a
thread. We can assess the accuracy of assembly sequences by analysing their thread
representation in the adjacency graph. Let $P$ be the thread representing an assembled sequence
in $G$. Any edge $e$ in $P$ is consistent if that edge is also labelled with segments from either or both
of the haplotypes. For any $P$ a contig path is a maximal subpath of $P$ in which all the edges are
consistent. Thus $P$ can be divided up into a series of contig paths, possibly interspersed with
edges in $P$ that are not contained in a contig path, see Figure 3 for an example. The bp length of
a contig path is equal to the sum of the bp lengths of the block edges it contains. Contig paths
represent maximal portions of the assembled sequence that are consistent with one or both of the
haplotypes and contain no assembly gaps, they can be thought of as portions of an assemblies’
contigs that perfectly follow a path through the graph of haplotype polymorphism.
Figure 2 shows contig path NG50s, defined analogously to block NG50; Supplementary Figures 12 and 13 show contig path coverage across the haplotypes, while Supplementary Figures 14 and 15 show in contrast the same plots, but instead use raw contig lengths. The contig path NG50s are substantially larger than block NG50s, for example the BGI assembly has a contig path NG50 1.5 orders of magnitude bigger than its block NG50. The difference between the largest and smallest block NG50 is 2,556 bp (GACWT 1,351 bp to BGI 3,907 bp); the difference between the largest and smallest contig path NG50 is 79,731 bp (GACWT 2,533 bp to BGI 82,264 bp). Thus the contig path NG50 results demonstrate that assemblies are able to reconstruct substantial regions perfectly, and contig path NG50 appears to be a more discriminative statistic than block NG50, as it indicates large differences between the assemblies.

3.4.5 Scaffold paths

To account for gaps within scaffolds, which we henceforth call scaffold breaks, we define scaffold paths. Scaffold paths can be thought of as portions of the assemblies’ scaffolds that perfectly follow a path through the graph of haplotype polymorphism, but which are allowed to jump unassembled sequences at scaffold gaps. Scaffold gaps are scaffold breaks (denoted as contiguous runs of wildcard characters in an assembly) whose surrounding contig ends are bridged by a path of haplotype representing edges within the adjacency graph, see Figure 3 for an example and the methods for a formal definition.

Notably, our definition of a scaffold gap within the graph is permissive in that it allows (1) any sequence of Ns to define a scaffold break and (2) the sequence of Ns that define the scaffold break to be aligned within the ends of the block that sandwich the gap in the assembly. This definition was sought because there is currently wide variation in the syntax used to define such gaps within different assemblers and to be tolerant of alignment errors caused by the phenomena of edge wander (Holmes & Durbin, 1998), caused when the alignment of positions around a gap has more than one equally probable scenario. As a scaffold path is a concatenation of contig paths its bp length is just the sum of the bp lengths of the contig paths that it contains.

Figure 2 shows the scaffold path NG50, defined analogously to the block and contig path NG50s, sorted with respect the scaffold NG50. In many cases the scaffold path NG50 is substantially larger than the contig path NG50. Figure 4 shows a stack fill plot of the coverage of scaffold paths along the three chromosomes of haplotype $\alpha$, (see also Supplementary Figures 16-19). It demonstrates the substantial differences between the assemblies and that large, megabase regions of the haplotype can be reconstructed with assembly gaps, but without apparent error.
3.4.6 Structural errors

Despite the long lengths of many scaffold paths, for most assemblies the scaffold NG50 is substantially larger than the scaffold path NG50, indicating that there were apparent errors that broke scaffolds into smaller sets of scaffold paths. To analyse these errors we continued our graph analysis, defining a number of subgraph types to represent them, which we formally define in the methods. These subgraph definitions include erroneous intra and inter chromosomal joins, insertions, deletions, simultaneous insertion and deletions and insertions at the ends of assembled sequences. Table 5 (and Supplementary Table 3) shows the numbers of structural errors for each assembly; Supplementary Figures 20 and 21 show structural errors across the haplotypes. Many assemblies do not have categories of error to which they are particularly prone, but a few do. In these cases there may be a systematic bias in the operation of the programs that generated them or in the way that we interpreted them.

Insertion and deletion (indel) structural errors involve, respectively, the addition and removal of contiguous run of bases. In Supplementary Figures 22 and 23 we investigate the size distribution of such errors, using both the described MSA and supporting alignments from the progressiveMauve program (Darling, Mau, & Perna, 2010). We find that in almost all cases the size distribution of the segments of inserted and deleted bases follows an approximately geometrically decreasing distribution.

We also searched for subgraphs in which the assembly created a haplotype to contamination chimera, but did not find any such subgraphs. To investigate this surprising result we searched for such chimeras using BLAST with relaxed parameters (Supplementary Section 7.1.3 and Supplementary Figure 24). Using this approach we found 56 assemblies with no such chimeras, 7 assemblies with 1 chimera, 1 assembly with 2 chimeras and one outlier assembly with 26 chimeras. In each case we verified that these chimeras were missed in the graph approach due to the stringent MSA parameters.

3.4.7 Long-range contiguity

The MSA graph theoretic analysis we have described is local in nature and quite strict, in that it has no notion of large-scale contiguity, and refuses to stitch together paths that would be joined, but for a small error. We thus sought a method to analyse the larger scale contiguity between pairs of separated points in the genome. Formally, for two positions $x_i$ and $x_j$ in a haplotype
chromosome $x$, such that $i < j$, if there exists two positions $y_k$, $y_l$ in an assembly scaffold $y$ such that (1) $y_k$ is in the same column as $x_i$, (2) $y_l$ is in the same column as $x_j$ and (3) $k < l$, we say $y_k$ and $y_l$ are correctly contiguous. Pairs may be correctly contiguous but not necessarily covered by the same contig path or scaffold path, and indeed there may be arbitrary numbers of assembly errors between two correctly contiguous positions.

Figure 5 shows the proportion of correctly contiguous pairs as a function of the pairs’ separation distance for each assembly. Taken at a high level, in all assemblies the proportion of correctly linked pairs monotonically decreases with separation distance. Therefore we take the separation distance at the 50th percentile, termed the Correct Contiguity 50 (CC50) as an essentially sufficient statistic. See Supplementary section 7.1.2 and Supplementary Figures 25 and 26 for supporting BLAST based analysis.

3.4.8 Annotation analysis

Evolver maintains annotations for a number of classes of simulated sequence, including genes, which Evolver models as having exons, introns and untranslated regions (UTRs), and conserved non-coding elements. Additionally, while Evolver does not track the history of individual repeat elements following their insertion, it maintains a library of mobile elements, and thus, using RepeatMasker (v1.25 http://www.repeatmasker.org) and tandem repeats finder (v4.0, http://tandem.bu.edu/trf/trf.html) with this library, we identified a subset of repetitive sequence within the haplotypes. Continuing the previous MSA analysis, we define a perfect path as a maximal subpath of a haplotype thread that is isomorphic to a subpath of an assembly thread. For a given assembly the corresponding set of perfect paths reflects the regions of the haplotypes that are perfectly reconstructed. Unlike contig and scaffold paths, perfect paths are intolerant of haplotype polymorphism, but give a well defined set of intervals within $\alpha_{1,2}$ for comparison with a set of annotations. Table 6 shows for each assembly the proportion of each annotation type contained within perfect paths.

Both haplotypes of the $\alpha_{1,2}$ genome contain 176 protein-coding genes, Supplementary Figure 27 show the distribution of their lengths; we find that only a small proportion (max 11% of bp, min 2% of bp) of these full length transcripts are perfectly reconstructed by the assemblies. Conversely, we find that in the best assemblies almost all exons and a high proportion of UTRs are perfectly reconstructed, for example, 99% of bp in exons and 84% of bp in UTRs of the BGI assembly. We
also find that most perfectly reconstructed genes are intron-less (data not shown), the assemblies therefore fail mostly to reconstruct introns perfectly. To further characterise this failure we used tBLASTN (Altschul, Gish, Miller, Myers, & Lipman, 1990) (see Supplementary Section 7.1.4) to align the spliced transcripts of $\alpha$ (without introns and UTRs) to the scaffolds of the assemblies, counting a match if it included 95% of the given transcript, see last column of Table 4 labelled genic correctness and Supplementary Table 1. This more tolerant analysis reveals that in the best assemblies the majority of exon chains are reconstructed contiguously (in the correct order and orientation) within single scaffolds, e.g. the Broad assembly has 107 spliced transcripts (93.8% of bp) reconstructed by this metric.

As expected, repeats were the least well reconstructed annotation types, with the best assembly, BGI, reconstructing only 64% of repeats perfectly (see last column of Table 6). As these regions are naturally difficult to align correctly within an MSA, we also performed BLAST based fragment analysis, see Supplementary Section 7.1.5, with similar results.

Finally, we also looked at conserved non-coding regulatory elements, which Evolver both models and tracks. As these elements are short and relatively non-repetitive the majority (88% - 99% of bp) were perfectly reconstructed by the assemblies.

### 3.4.9 Substitution errors

We have so far described assessments of structural correctness and contiguity, both overall and for functional genic elements. We now turn to the assessment of somewhat orthogonal issues, firstly by looking at base calling and then finally by analysing copy number errors.

Although we do not allow structural rearrangements within MSA blocks, blocks are tolerant of substitutions. Let a (haplotype) column of aligned bases within a block that (1) contains a single position from both haplotypes and (2) a single position from an assembly sequence be called valid. We use these criteria because such columns unambiguously map a single assembled sequence to a single position in the alignment of both haplotypes, while avoiding the issues of paralogous alignment and multiple counting. We distinguish two types of valid columns: (1) homozygous columns, those containing the same base pair from both haplotypes and (2) heterozygous columns, those containing distinct base pairs from each haplotype. We also initially considered columns that contain one but not both haplotypes, but found that the numbers of such columns that we could consider reliably aligned was not sufficient for us to confidently compute statistics.
Assemblers are free to use IUPAC ambiguity characters to call bases. To allow for this we use a bit-score to score correct but ambiguous matches within valid columns (see methods). We say there has been a substitution error if the position in the assembly sequence has an IUPAC character that does not represent either of the haplotypes’ base pair(s).

Some of the substitution errors that we observe are likely due to misalignments. These can occur due to edge wander (Holmes & Durbin, 1998), or the larger scale misalignment of an assembled sequence to a paralog of its true ortholog. The sum of substitution errors over all valid columns is therefore an upper bound on the substitution errors within valid columns. To obtain a higher confidence set of substitution errors we select a subset of valid columns that meet the following requirements: (1) Are part of blocks of at least 1 kb in length, avoiding errors within short indels, (2) are not within 5 positions of the start and end of the block, avoiding edge wander, and (3) are within blocks with 98% or higher sequence identity, ensuring the alignments are unlikely to be paralogous. The sum of substitution errors within these high confidence valid columns represents a reasonable lower bound of the number of substitution errors within valid columns.

Figure 6 (also Supplementary Figure 29 and Supplementary Tables 4 and 5) show, as might be expected, that there are, in general, proportionally more errors made in heterozygous columns than homozygous columns, though there are naturally much fewer overall heterozygous positions, for example the WTSI-S and CRACS assemblies made no heterozygous errors. We find a strong correlation between error rates in heterozygous and homozygous columns, with the exceptions of the Broad and BCCGSC assemblies, which have proportionally higher rates of heterozygous errors. The Broad result is explained by the large number of N ambiguity characters called at heterozygous sites, which makes the number of errors per bit correspondingly higher, while the BCCGSC result was due to a programmatic error in the assembler’s pipeline that has since been identified and resolved as a result of this analysis. Interestingly, we find considerable variation between the programs in overall error rates. The strongest assembly, WTSI-S, makes one error for every $15.3 \times 10^6$ to $2.94 \times 10^6$ correct bits, or approximately one every $7.7 \times 10^5$ to $1.49 \times 10^5$ bases, while the weakest assembly, UCSF, makes an error for every $6.7 \times 10^3$ to $1.81 \times 10^4$ correct bits, or approximately one in every $3.3 \times 10^3$ to $9.0 \times 10^3$ bases.

### 3.4.10 Copy-number errors

Within any haplotype column of the MSA the copy number of the simulated diploid genome can be described by an interval $[\text{min}, \text{max}]$, where min is the minimum number of bases either of the
two haplotypes contributes and max is the maximum number of bases either of the two haplotype contributes. To establish if assemblies were producing too many or too few copies of the homologous positions within the two haplotypes we looked for haplotype columns where the copy number of the assembly lay outside of this copy number interval. There are two possibilities, either the number of copies in the assembly is less than min, in which case there is a deficiency in the copy number, or the number of copies in the assembly is greater than max, in which case there is an excess in the copy number.

Figure 7 (also Supplementary Figure 30) shows the proportions of haplotype columns with copy number excesses and deficiencies. Again, to address contributions made by alignment errors we choose to produce an upper and lower bound on these proportions. The lower bound is taken over all haplotype columns in the alignments, while the upper bound is computed over only haplotype columns that are part of blocks of at least 1 kb in length.

The deficiencies are dominated by cases where the assembly is not present; therefore copy number deficiency is closely correlated with coverage. Unfortunately, there are not a sufficient number of cases where the assembly is present but the copy number is deficient so that we may make reliable inferences about this interesting category. This appears to be a consequence of the genome simulation lacking sufficient numbers of recent duplications, and may be an indication that the genome simulation is somewhat unrealistic, as other authors (Worley & Gibbs, 2010) (Alkan, Sajjadian, & Eichler, 2011) have discussed that recent segmental duplications cause substantial problems for assemblies generated with short reads.

We find that there are substantial numbers of copy number excesses, such that generally the number of excesses was larger than the number of deficiencies. We find that excesses do not correlate particularly well with deficiencies, particularly for programs with extremes of deficiency or excess. We do find, however, that excesses correlate well with input assembly size (data not shown). The best assembly, EBI, has excesses in between 0.0521% and 0.752% of haplotype columns, while the least assembly, nABySS has excesses in between 30.8% and 33.5% of haplotype columns.

4. Discussion

We have used simulation to create a novel benchmark dataset for de novo assembly. We have evaluated a previously unprecedented 41 different assemblies from 17 different groups, making it the largest short read de novo assembly evaluation to date. In summary, we have assessed
coverage, the lengths of consistent contig and scaffold paths, structural errors, long-range contiguity, the assembly of specific annotated regions, including genes and repeats, base calling errors and copy number errors; Table 7 conveniently summarises these evaluation metrics. This benchmark dataset is freely available online at http://www.assemblathon.org/ and is supplemented by code that can take new assemblies and amalgamate the new result into the analysis we present here. It is our hope that this standard will assist the assembly community when introducing new methods by providing a large set of metrics and methods with which to compare.

Given the degree of polymorphism within the α1,2 genome the haplotype aware evaluations proved critical to the assessment. For example, the haplotype aware path analysis demonstrates that methods are able to reconstruct multiple megabases, with scaffold breaks, essentially perfectly. We chose to treat switches between the haplotypes of α1,2 permissively because the assemblers were not asked to reconstruct the two haplotypes, but rather to produce a consensus reference of the two. It is an open question if, with this dataset or one like it, an assembly could produce phased variants of each scaffold. In Section 7.3 of the supplement we test if there was any evidence of teams phasing single nucleotide polymorphisms (SNPs) or structural variants by preferentially choosing one haplotype, but we do not find convincing evidence for either, apart from that inadvertently caused by a bias in the simulated reads (see Section 5.2.5).

Table 3 shows the rankings of each of the featured assemblies for each of the described assessments; additionally in Supplementary Figures 31 and 32 we assess correlations between the logs of different metrics. Intuitively, one might expect the path analysis metrics and the contiguity assessments to be correlated to one another and inversely correlated with structural errors. Indeed, this intuition proves partially correct. Contiguity (CC50) and scaffold path NG50 are strongly correlated (R² = 0.77, P < 0.001), while structural errors are inversely correlated with scaffold path NG50s, with one explaining about half the variance of the other (R² = 0.48, P < 0.001). However, contig path NG50 is only weakly correlated with scaffold path NG50 (CPNG50 – SPNG50 R² = 0.38, P < 0.001) and contiguity (CPNG50 - CC50 R² = 0.31, P < 0.01), suggesting that the scaffolding process is more important in producing accurate long scaffolds than the prior contigging process.

Given the popularity and simplicity of N50 statistics, it is perhaps reassuring how well these metrics correlate with the path and contiguity metrics (SN50 – CC50 R² = 0.98, P < 0.001; SN50 –
SPNG50 $R^2 = 0.74$, $P < 0.001$; CN50 – CPNG50 $R^2 = 0.64$, $P < 0.001$), suggesting that one may usefully compare N50 measurements between assemblies, and not just between assemblies by the same program. Interestingly, the genic correctness measure also correlates with all the N50 measures, and most strongly with the correct contiguity 50 ($R^2 = 0.98$, $P < 0.001$) and scaffold N50 measures ($R^2 = 0.98$, $P < 0.001$).

We do not find that substitution errors and copy number errors correlate substantially with anything else, except for a correlation between substitution errors and structural errors ($R^2 = 0.45$, $P < 0.001$). This is perhaps unsurprising, given the orthogonal basis of these metrics to each other and the other evaluations. Perhaps surprisingly, coverage does not correlate strongly with other measures, and in particular not with contig or scaffold N50 statistics, suggesting such naïve measures are not good proxies for coverage.

Table 3 highlights that while the best assemblies are stronger in most categories than the weakest assemblies, all the assemblies have areas in which they can improve relative to their peers, if at a trade-off cost in other categories. For example, the BGI assembly, while having the largest contig path NG50, has only the sixth largest scaffold path NG50, which is more than 4 times smaller than the strongest method in this category (WTSL-S), suggesting that its scaffolding could be improved. Conversely, the WTSI-S and DOEJGI assemblies had large contiguity (CC50) and scaffold path (SPNG50) measures and low numbers of structural errors, but relatively short contig paths (CPNG50), suggesting their contigging could be made more aggressive, though possibly with a corresponding increase in structural errors.

We have demonstrated in simulation that the best current sequence assemblers can reconstruct at high coverage and with good accuracy large sequences of a substantial de novo genome. This is concordant with other recent work that suggests that short read sequencing is becoming competitive with capillary sequencing (Maccallum, et al., 2009) (Gnerre, et al., 2011). MacCallum et al looked at five microbial genomes with sizes ranging from 2.8 Mb (Staphylococcus aureus) to 39.2 Mb (Neurospora crassa) and determined that with data from two paired libraries that the ALLPATHS 2 program was able to produce assemblies with qualities that exceeded draft assemblies using Sanger methods. Gnerre et al sequenced two genomes; a human cell line (GM12878) and a mouse strain (C57BL/6J female) and assembled them using the ALLPATHS-LG program. The authors found that with Illumina reads of 100x coverage in four library types that
their assemblies neared capillary sequencing quality in completeness, long range connectivity, contiguity and accuracy.

There are a number of important limitations with the current work. Firstly, the use of simulation makes it hard to know how applicable these results are to any other dataset; though this is arguably true of any dataset, the simulation’s limitations, in particular the noted issues with the read simulation and with the low repeat content of the genome, likely influence the results. Secondly, the limited size of the simulated genome means that some of the strategies employed here may not work as effectively, or at all, on larger vertebrate scale genome datasets. Finally, as our results derive from a single dataset in which no attempt was made to measure the variance of our various metrics, it is questionable how reliable our measurements are. To address these issues a second competitive evaluation, Assemblathon 2, is now underway.

Given the scale of challenges in making assessments, and to avoid fragmentation, we suggest that Assemblathon 2 continue to focus on the assessment of complete pipelines, rather than attempting to assess individual pipeline components, and that it continue to rely on the individual assembly teams to compute their own assemblies, despite this making it difficult to compare the computational requirements of the pipelines, given the self reported nature of such data and the heterogeneous equipment upon which the assemblies are computed.

However, we conclude by making three distinguishing suggestions for Assemblathon 2 that would sufficiently expand its scope from this initial competition. Firstly, it should feature at least one mammalian genome scale data set, to test the scaling of the assembly pipelines. Secondly, it should feature real data, to compare with the simulation results presented in this competition; this may necessitate the use of a different set of evaluation metrics, where the “correct” answer is unknown. Thirdly, it should be expanded to include other sequencing technologies, so that a better comparative, unbiased understanding of available sequencing technologies can be made.

5. Methods

5.1 Genome simulation

The Evolver simulation was managed by a set of scripts (https://github.com/dentearl/evolverSimControl/), which control the execution of Evolver and allow a general phylogeny to be simulated.
As well as a starting sequence, Evolver also requires a set of annotations that are used assign sequences to element types that undergo differential evolution simulation. The following annotations were used: UCSC Genes, UCSC Old Genes, CpG Islands, Ensembl Genes and MGC Genes from the UCSC table browser (Fujita, et al., 2011). The root genome was then coupled with parameters and a mobile element library provided by Arend Sidow (pers. comm.) to form the Evolver in-file set for the simulation.

Evolver proceeds iteratively by a series of discrete steps. We used an Evolver step length of 0.01 substitutions per site, meaning the initial branch length of 0.4 substitutions per site (~200 my) (Fujita, et al., 2011) (Hedges, Dudley, & Kumar, 2006)) from the root node to the most recent common ancestor (MRCA) of the final leaf genomes node consisted of 40 separate Evolver cycles. The lineages leading from the MRCA to $\alpha$ and $\beta$ descend for a distance of 0.1 (~50 my) substitutions per site in 10 Evolver cycles. The final splits into the lineages leading to the leaf genomes were each performed in 1 Evolver cycle of 0.002 substitutions per site (~1 my), with parameters scaled appropriately. An alignment between the $\alpha_1$ and $\alpha_2$ haplotypes is available on the project website (http://compbio.soe.ucsc.edu/assemblathon1/).

### 5.2 Read Simulation

Prior to writing our own read simulator we considered several pre-existing tools. We first considered wgsim (Li, et al., 2010). Unfortunately, this program does not model mate-pair Illumina reads, and it models error rates uniformly across the sequence. We note that this error rate limitation is removed in dwgsim (http://sourceforge.net/apps/mediawiki/dnaa/). However, dwgsim does not model chimeric mate-pair reads or paired-end contamination, which we wished to model. We contacted Illumina and requested their in-house programs for simulating reads. The Illumina software package was capable of modelling chimeric mate-pair reads, and it modelled error rates by copying quality strings from a user supplied file of Illumina reads. Unfortunately, this method did not allow us to model different error rates conditioned on different underlying bases, which we felt was important. We also considered several other software packages for modelling Illumina style reads, including metasim (Richter, Ott, Auch, Schmid, & Huson, 2008), PEMer (Korbel, et al., 2009), ReSeqSim (Du, Bjornson, Zhang, Kong, Snyder, & Gerstein, 2009), SimNext (http://evolution.sysu.edu.cn/english/software/simnext.htm), Flux Simulator (http://flux.sammeth.net/index.html), and Mason (part of the SeqAn package (Döring, Weese, Rausch, & Reinert, 2008)), all of which lack one or more of the criteria we desired.
Given these findings we wrote our own simulator, which combined the capabilities of the Illumina supplied software to model chimeric mate-pair reads as well as standard pared-end reads, with our own position and reference-base-specific empirical error model trained on Illumina data.

5.2.1 Read Sampling Strategy

For read sampling we employed two separate methods, one for mate-pair libraries and the other for paired end libraries. Reads were first sampled uniformly across each sequence. Coverage depth was kept approximately uniform by weighting the number of reads sampled from each sequence by its length. Read fragments were sampled from either strand with equal probability. Duplicates were produced with some probability before the error was applied to the reads. See Supplementary Figure 33 for a density map of read depth across the haplotypes.

5.2.2 Paired-End Sampling

Illumina paired end sampling was the most straightforward strategy to simulate. It involved randomly selecting fragments in the 150-500 base pair range uniformly across the genome until the desired coverage was met (specific sizes below). Fragment size was sampled from a normal distribution with a specified mean and variance. The reads were oriented facing each other and were sampled from either strand with equal probability. The following paired-end libraries were generated:

- 200 bp insert +/- 20 standard deviation
  - 2x 100 bp
  - 22,499,731 read pairs (~40x coverage of the diploid sequence)
  - 0.01 probability of being a duplicate
- 300 bp insert +/- 30 standard deviation.
  - 2x 100 bp
  - 22,499,731 read pairs (~40x coverage)
  - 0.01 probability of being a duplicate

5.2.3 Mate-Pair Sampling

Illumina mate-pair library construction differs from paired-end library construction in that it introduces several unique types of error into the reads. In reality these libraries are constructed by attaching a chemical tag onto the ends of a long sequence fragment, typically in the range of 2-10 kb, after which the fragment is circularized. The circularised product is then further fragmented.
into sizes typically within the 200-500 bp range, which is the upper limit on fragment lengths for Illumina sequencing. Finally the resulting mixture is purified for fragments that contain the chemical tag, so that DNA from near the ends of the original 2-10 kb loop are what ideally get sequenced.

There are three common types of error introduced in the mate-pair library preparation process, and we modelled two of them. Firstly, when the fragments are circularized, there is a chance that a loop will be formed between two non-related long fragments, resulting in chimeric reads between two unrelated parts of the genome. We did not model this type of error. Assuming that the fragment is properly circularised, the second type error is produced when a fragment that does not contain the chemical tag is mistakenly sampled. When this happens, the loop join is not part of the fragment, and a paired-end style read with a short insert is mixed in with the rest of the library. We did model this type of error, and varied the probability of its occurrence with each mate-pair library. The final major source of error is created during the random fragmentation process and results in the loop join position occurring in the middle of a read rather than between the two reads. We modelled this by assuming a uniform distribution of loop join sites across a sampled loop fragment, which resulted in chimeric reads as a function of the size of the fragmented loop piece, and the length of the reads. For example shorter reads and longer loop fragmentation pieces were less likely to result in a chimeric read. The following mate-pair libraries were generated:

- 3 kb loop length +/- 300 standard deviation
  - 2x 100 bp
  - 500 bp loop fragmentation size +/- 50 bp
  - 0.2 probability of sampling a PE fragment rather than an MP fragment
  - 11,249,866 read pairs (~20x coverage)
  - 0.05 probability of being a duplicate

- 10 kb loop length +/- 1 kb standard deviation
  - 2x 100 bp
  - 500 bp loop fragmentation size +/- 50 bp
  - 0.3 probability of sampling a PE fragment rather than an MP fragment
  - 11,249,866 read pairs (~20x coverage)
  - 0.08 probability of being a duplicate

5.2.4 Base-Level Error Model
We utilized an error model that is dependent on the position within the read and the underlying reference base. To generate this model we assembled a human mitochondrial genome using reads from an Illumina HiSeq run (http://www.illumina.com/systems/hiseq_2000.ilmn) with the reference guided assembler MIA (Green, et al., 2010). We then took that assembly and mapped all reads back to it using BWA with default settings, to do a paired end mapping to the sequence. We kept all alignments with a mapq quality score over 10. We then iterated through the alignment and built an empirical distribution of Phred (Ewing, Hillier, Wendl, & Green, 1998) scores and the probabilities of observing one of A, C, G, T or N given the reference base, the position in the read, and the reported Phred quality score. The error model was therefore conditioned on the Phred score, position, and reference base, and did not assume that the Phred scores were an accurate representation of the underlying error rates.

5.2.5 Problems With the Error Model Used in Assemblathon 1

The error model used was appealingly simple but has limitations that should be understood. Firstly, in generating the error model we omitted many reads that had an error rate that was too high to confidently map to the assembled mitochondria. In the future this could partially be overcome by using the PhiX control lane (http://www.illumina.com/products/multiplexing_sequencing_primers_and_phix_control_kit.ilmn), where one can confidently force the vast majority of the reads to map back to the PhiX 174 genome (NCBI accession NC_001422.1), and do not have to be as sensitive to false positive alignments.

Secondly, since we used a flat naive prior on the distribution of Phred scores, when training our empirical model there was, due to noise, a mixture of good and poor quality bases at the ends of the reads. Since each position was treated independently, the distribution of Phred scores was therefore likely not typical, resulting in the likely relative failure of assembler heuristics used to trim strings of bad Phred scores at the ends of reads.

Thirdly, since we wrote the simulator following the general algorithmic flow of the wgsim read simulator (Li, et al., 2010), reads were randomised within haplotype chromosomes, but not between haplotype chromosomes, resulting in reads from each haplotype and chromosome being clustered together separately in the data. Thankfully, an investigation of phasing bias in Supplementary Section 7.3 shows only a couple of assemblies showed evidence of any bias that could likely be attributable to this.
5.3 Cactus Alignment Assessment

5.3.1 Alignment Generation

The Cactus program starts by using the Lastz pairwise alignment program (http://www.bx.psu.edu/~rsharris/lastz/) to generate a set of pairwise alignments between all the input sequences, including intra-sequence alignments that arise from recent duplications. In the adapted version of Cactus used for the Assemblathon, which we henceforth call Cactus-A, we used the following parameters to Lastz, after discussion with the program's author: --step=10 --seed=match12 --notransition --mismatch=2,100 --match=1,5 --ambiguous=iupac --nogapped --identity=98. This ensured that the resulting pairwise alignments were ungapped (without indels), of minimum length of 100 bp and with an identity (sequence similarity) of 98% or greater, in concordance with the evolutionary distance between the haplotypes. Cactus-A uses these alignments to build a “sparse map” of the homologies between a set of input sequences. Once this sparse map is constructed, in the form of a Cactus graph (Paten, et al., 2011), a novel algorithm is used to align together sequences that were initially unaligned in the sparse map. To prevent sequences that are not homologous from being aligned in this process we set the alignment rejection parameter, called $\gamma$, to 0.2, to filter positions from being aligned that are not likely to have very recently been diverged. The results of Cactus-A are stored as MAF files (Blanchette, et al., 2004), one for each assembly, these are available in the supplementary material.

5.3.2 Scaffold Gaps, Error Subgraphs and Scaffold Paths

Let $P$ be a sequence of block edges $((x_1, x_2), (x_3, x_4) \ldots (x_{n-1}, x_n))$ in a thread (thus ignoring the alternating adjacency edges $(x_2, x_3), (x_4, x_5)$ etc.) representing an assembled sequence in the adjacency graph. The ambiguity of a sequence is equal to the number of wildcard characters that it contains (denoted as $N_s$). Similarly, the ambiguity of a subsequence of $P$ is equal to the ambiguity of the subsequence of the assembly sequence it represents. The prefix ambiguity of $(x_i, x_j)$ is equal to the number of wildcard characters in the first 5 bases of the assembly sequence that $(x_i, x_j)$ represents, orienting the sequence from $x_i$ to $x_j$. The approximate ambiguity of a subsequence $Q = ((x_1, x_{i+1}), (x_{i+2}, x_{i+3}) \ldots (x_{n-1}, x_n))$ is equal to the ambiguity of $Q$ plus the prefix ambiguity of $(x_i, x_{i+1})$ and $(x_{i+j-1}, x_{i+j+2})$, if these edges exist. By using approximate ambiguity rather than just ambiguity we allow for wobble in the alignment caused by edge wander (Holmes & Durbin, 1998) when denoting a scaffold gap.
We say a thread is *empty* if it represents a sequence of zero length, else we say it is *non-empty*.

Let a maximal thread of inconsistent adjacency edges and block edges that do not contain haplotypes or bacterial contamination segments be called a *joining thread*. A joining thread represents an unaligned portion of an assembly sequence. A scaffold gap or error subgraph is defined by a joining thread incident at one or both ends with blocks that contain haplotype segments. We classify such joining threads as follows:

(A) If the joining thread is not attached to anything at one end (i.e. it terminates) (Figure 8A):

- If it has approximate ambiguity then we classify it as a scaffold gap.
- Else we classify it as a hanging insert error.

(B) If the joining thread is attached at each end to blocks, $a$ and $b$, containing haplotype segments:

- If $a$ and $b$ are connected by a thread containing haplotype segments (Figure 8B):
  
  i. If the joining thread has approximate ambiguity then it is a scaffold gap.
  
  ii. Else it is an indel (insertion/deletion) error:

  1. If the joining thread is empty then it is a deletion error, by definition all haplotype paths between $a$ and $b$ must be non-empty.
  
  2. Else if all haplotype threads are empty then it is an insertion error, by definition the assembly thread must be non-empty.
  
  3. Else, all the haplotype threads and the assembly thread are non-empty and it is an insertion and deletion error.

- Else $a$ and $b$ are not attached by a thread of haplotype containing edges:
i. If \( a \) and \( b \) both contain positions from one or more common haplotype sequences then it is an intra haplotype joining error (Figure 8C).

ii. Else it is an intra haplotype joining error (Figure 8C).

(C) Else, the joining thread is attached at one end to a bacterial contamination containing block (Figure 8B) and we classify it as a haplotype to contamination joining error (Figure 8C).

For any thread \( P \) representing an assembly sequence, a scaffold path is a maximal subpath of \( P \) in which all the edges are consistent and/or part of a scaffold gap subgraph.

5.3.3 Substitution errors

We use a bit-score to score correct but ambiguous matches within valid columns. We assign to each valid column the bit score \(-m \log_2(n/4)\), where \( n \) is the number of different bases the IUPAC character in the assembly represents and \( m \) is the number of distinct base pairs in the two haplotypes that matches or is represented (amongst others) by the assembly IUPAC character. Thus in homozygous columns the score is at most 2, in heterozygous columns the score is 2 if and only if the assembly correctly predicts one of the two base pairs, or if it predicts an ambiguity character that represents both and only those two base pairs.

6. Acknowledgements

We would like to thank Bob Edgar, Arend Sidow and George Asimenos for their help with using Evolver. We thank three anonymous reviewers for comments and discussion on previous versions of this manuscript. We acknowledge the following grants: ENCODE DAC (data analysis center) subaward on NHGRI grant no. U01HG004695 to the European Bioinformatics Institute; ENCODE DCC (data coordination center) NHGRI grant no. U41HG004568; Browser (Center for Genomic Science) NHGRI grant no. P41HG002371; Gencode subaward on NHGRI grant no. U54HG004555 to the Sanger Center; NCI 1U24CA143858-01; NIH HG00064; PTDC/BIA-BEC/100616/2008; PTDC/EIA-EIA/100897/2008; the Fundacao para a Ciencia e Tecnologia; National Natural Science Foundation of China (30725008; 30890032; 30811130531; 30221004); a National Basic Research Program of China (973 program no. 2011CB809200); and the Chinese 863 program (2006AA02Z177; 2006AA02Z334; 2006AA02A302; 2009AA022707).


Figure 1 — The phylogeny of the simulated haploid genomes. The root genome derives from human chromosome 13. The $\alpha_1$ and $\alpha_2$ haplotypes form the diploid genome from which we generated reads. The $\beta_1$ and $\beta_2$ haplotypes form a diploid out-group genome that was made available to the assemblers.
Figure 2 — N50 statistics. Assemblies are sorted left to right in descending order by scaffold path NG50. Data points for each assembly are slightly offset along the x-axis in order to show overlaps.

Figure 3 — An adjacency graph example demonstrating threads, contig paths and scaffold paths. Each stack of boxes represents a block edge. The nodes of the graph are represented by the left and right ends of the stacked boxes. The adjacency edges are groups of lines that connect the ends of the stacked boxes. Threads are represented inset within the graph as alternating connected boxes and coloured lines. There are three threads shown: black, gray and red. The black and gray threads represent two haplotypes, there are many alternative haplotype threads that result from a mixture of these haplotype segments which are equally plausible given no
additional information to de-convolve them. The red thread represents an assembly sequence. For the assembly thread consistent adjacencies are shown in solid red. The dashed red line between the right end of block $g$ and the left end of block $i$ represents a structural error (deletion). The dashed red line between the right end of block $k$ and the left end of block $m$ represents a scaffold gap, because the segment of the assembly in block $n$ contains wild card characters. The example therefore contains three contig paths: (from left to right) blocks $a...g$ ACTGAAATCGGGACCCC; blocks $i, j, k$ GGAAC; and block $m$ CC. However the example contains only two scaffold paths because the latter two contig paths are concatenated to form one scaffold path.

Figure 4 — Assembly coverage along haplotype $\alpha_1$ stratified by scaffold path length weighted overall coverage. The top 6 rows show density plots of annotations. CDS: coding sequence; UTR: untranslated region; NXE: non-exonic conserved regions within genes; NGE: non-genic conserved regions; island: CpG islands; repeats: repetitive elements. The remaining rows show the top ranked assembly from each group, sorted by scaffold path length weighted overall coverage. Each such row is a density plot of the coverage, with coloured stack fills used to show
the length of scaffold paths mapped to a given location in the haplotype. For example, the left most light-orange block of the WTSI-S assembly row represents a region of haplotype \( \alpha \), that is almost completely covered by a scaffold path from the WTSI-S assembly greater than one megabase in length.

Figure 5 — The proportion of correctly contiguous pairs as a function of their separation distance. Each line represents the top assembly from each team. Correctly contiguous 50 (CC50) values are the lowest point of each line. The legend is ordered top to bottom in descending order of CC50. Proportions were calculated by taking 100 million random samples and binning them into 2,000 bins, equally spaced along a \( \log_{10} \) scale, so that an approximately equal number of samples fell in each bin.
Figure 6 — Substitution (base) errors for the top assembly from each team. Top: substitution errors per correct bit within all valid columns, middle: substitution errors per correct bit within homozygous columns only, bottom: substitution errors per correct bit within heterozygous columns only. Assemblies are sorted from left to right in ascending order by the sum of substitutions per correct bit. In each facetted plot each assembly is shown as an interval, giving the upper and lower bounds on the numbers of substitution errors (see main text).
Figure 7 — Copy number errors for the top assembly from each team. Top: proportion of haplotype containing columns with a copy number error, middle: proportion of haplotype containing columns with an excess copy number error, bottom: proportion of haplotype containing columns with an excess copy number error. Assemblies are sorted from left to right in ascending order according to the proportion of haplotype containing columns with a copy number error. In
each facetted plot each assembly is shown as an interval, giving the upper and lower bounds on the numbers of copy number errors (see main text).

Figure 8 — Scaffold gap and error subgraphs. Diagrams follow the format of Figure 3. The rounded boxes represent extensions to the surrounding threads. Line ends not incident with the edge of boxes represent the continuation of a thread unseen. In each diagram the right end of block a and the left end of block b (if present) represent the ends of contig paths, the enclosed red thread represents the joining thread. The black thread represents a haplotype thread. The gray thread represents either a haplotype or bacterial contamination thread. (A) Represents (hanging) scaffold gaps and hanging insert errors. (B) Represents scaffold gaps and indel errors. (C) Represents intra and inter chromosomal joining errors and haplotype to contamination joining errors.
Table 1 — Groups that submitted assemblies. The first 17 rows in the table correspond to entries submitted by participants in the competition. Assemblies with IDs beginning with “n,” (for naïve), were generated by organisers of the competition to demonstrate the performance of popular programs run with variations on their default parameters. *CSHL.1 used the β genome though that team’s top assembly, CSHL.2, which is referred to in the main paper as CSHL, did not.

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Table 2 — Genome simulation statistics. (A) Event numbers are between the previous branch point and the named node. Mb: size of the genome in megabases; GC: percentage GC content; Reps: percent of the genome masked by the union of tandem repeats finder and RepeatMasker, *is the published value for chromosome 13 (Dunham, et al., 2004); Reps 100mer: percent repetitiveness of the sequence and its reverse complement for 100-mers calculated with the
tallymer tool (Kurtz, Narechania, Stein, & Ware, 2008); Chr: number of chromosomes; Subs: number of substitution events; Dels: number of deletion events; Inv: number of inversion events; Moves: number of translocations; Copy: number of DNA segmental duplications; Tandem: number of tandem repeat insertions; Chr Split: number of chromosome fission events; Chr Fuse: number of chromosome fusion events. (B) Differences between haplotypes \( \alpha_1 \) and \( \alpha_2 \) as determined by inspection of the Evolver pairwise alignment. SNPs: count of single nucleotide polymorphisms; Subs: count of substitutions, including SNPs; \( \Sigma \) Subs: sum of the lengths of all substitutions; Indels: count of insertion deletion events; \( \Sigma \) Indels: sum of the lengths of all insertion deletion events; Inv: the sum of number of inversions invoked in each of the \( \alpha_1 \) and \( \alpha_2 \) Evolver steps.

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Table 3 — Rankings of the top assembly from each team in eight categories. For each category (listed below) all the received assemblies were ranked. The sum of the rankings from each category was then used to create an overall rank for the assemblies, the top (lowest number) ranked assembly from each group was then selected for inclusion in this manuscript. Numbers are ranks, with values shown in parentheses. Overall: sum of all rankings (possible range 8-160), CPNG50: Contig path NG50, SPNG50: Scaffold path NG50, Struct.: Sum of structural errors, CC50: length for which half of any two valid columns in the assembly are correct in order and orientation, Subs.: Total substitution errors per correct bit, Copy Num.: Proportion of columns with
a copy number error, Cov. Tot.: Overall Coverage, Cov. Genic: Coverage within coding sequences.

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Table 4 — Coverage statistics for the top assembly from each team. Hap Total: overall coverage, Hap $\alpha_1$: percent coverage for Haplotype $\alpha_1$, Hap $\alpha_2$: percent coverage for Haplotype $\alpha_2$, Bac: percent coverage of the bacterial contamination, Genic: percent coverage of the coding sequences, Unmapped: number of unmapped bases, many corresponding to short contigs.

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<th>Hap $\alpha_2$ (%)</th>
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<td>92.8</td>
<td>7.822e+05</td>
</tr>
<tr>
<td>WTSI-S</td>
<td>97.8</td>
<td>97.8</td>
<td>97.8</td>
<td>99.1</td>
<td>91.8</td>
<td>4.948e+06</td>
</tr>
<tr>
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<td>97.7</td>
<td>97.7</td>
<td>0.9</td>
<td>88.5</td>
<td>4.553e+05</td>
</tr>
<tr>
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<td>97.5</td>
<td>97.5</td>
<td>97.5</td>
<td>99.8</td>
<td>57.2</td>
<td>1.111e+07</td>
</tr>
<tr>
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<td>97.4</td>
<td>97.3</td>
<td>99.5</td>
<td>92.3</td>
<td>5.304e+06</td>
</tr>
<tr>
<td>nCLC</td>
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<td>97.2</td>
<td>97.2</td>
<td>99.8</td>
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<td>5.673e+06</td>
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<tr>
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<td>96.5</td>
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<td>84.8</td>
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<tr>
<td>CRACS</td>
<td>96.3</td>
<td>96.3</td>
<td>96.3</td>
<td>99.8</td>
<td>90.2</td>
<td>5.265e+06</td>
</tr>
<tr>
<td>DCISISU</td>
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<td>94.3</td>
<td>94.2</td>
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<tr>
<td>IRISA</td>
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<td>93.7</td>
<td>93.7</td>
<td>99.7</td>
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<tr>
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<td>90.9</td>
<td>90.9</td>
<td>100.0</td>
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<td>5.175e+06</td>
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<tr>
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<td>86.4</td>
<td>86.4</td>
<td>86.4</td>
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<td>48.0</td>
<td>2.053e+06</td>
</tr>
<tr>
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<td>83.7</td>
<td>83.7</td>
<td>83.7</td>
<td>0.0</td>
<td>59.6</td>
<td>1.822e+06</td>
</tr>
<tr>
<td>CIIUCc</td>
<td>78.5</td>
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<td>78.1</td>
<td>0.0</td>
<td>48.9</td>
<td>3.638e+06</td>
</tr>
</tbody>
</table>
Table 5 — Structural error statistics for the top assembly from each team. Columns are defined in the main text.

<table>
<thead>
<tr>
<th>ID</th>
<th>Intra chromosomal joins</th>
<th>Inter chromosomal joins</th>
<th>Insertions</th>
<th>Deletions</th>
<th>Insertion and deletion</th>
<th>Insertion at ends</th>
<th>Σ errors</th>
</tr>
</thead>
<tbody>
<tr>
<td>DOE/GI</td>
<td>21</td>
<td>160</td>
<td>55</td>
<td>108</td>
<td>40</td>
<td>72</td>
<td>456</td>
</tr>
<tr>
<td>WTDS-S</td>
<td>6</td>
<td>191</td>
<td>56</td>
<td>78</td>
<td>19</td>
<td>127</td>
<td>475</td>
</tr>
<tr>
<td>Broad</td>
<td>75</td>
<td>161</td>
<td>524</td>
<td>379</td>
<td>9</td>
<td>96</td>
<td>1,244</td>
</tr>
<tr>
<td>CRACCS</td>
<td>687</td>
<td>303</td>
<td>121</td>
<td>121</td>
<td>51</td>
<td>306</td>
<td>1,256</td>
</tr>
<tr>
<td>nABySS</td>
<td>17</td>
<td>48</td>
<td>208</td>
<td>188</td>
<td>63</td>
<td>1,207</td>
<td>1,731</td>
</tr>
<tr>
<td>BGI</td>
<td>368</td>
<td>288</td>
<td>355</td>
<td>639</td>
<td>98</td>
<td>130</td>
<td>1,878</td>
</tr>
<tr>
<td>EBI</td>
<td>458</td>
<td>563</td>
<td>127</td>
<td>547</td>
<td>53</td>
<td>307</td>
<td>2,055</td>
</tr>
<tr>
<td>RHUL</td>
<td>691</td>
<td>349</td>
<td>172</td>
<td>264</td>
<td>26</td>
<td>1,049</td>
<td>2,551</td>
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<tr>
<td>ASTR</td>
<td>2,065</td>
<td>200</td>
<td>109</td>
<td>227</td>
<td>73</td>
<td>144</td>
<td>2,818</td>
</tr>
<tr>
<td>BCCGSC</td>
<td>351</td>
<td>285</td>
<td>255</td>
<td>233</td>
<td>102</td>
<td>1,641</td>
<td>2,867</td>
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<tr>
<td>HRISA</td>
<td>147</td>
<td>203</td>
<td>925</td>
<td>1,593</td>
<td>136</td>
<td>741</td>
<td>3,725</td>
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<tr>
<td>DGSISU</td>
<td>1,410</td>
<td>956</td>
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<td>954</td>
<td>109</td>
<td>560</td>
<td>4,319</td>
</tr>
<tr>
<td>WTSI-P</td>
<td>1,940</td>
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<td>1,851</td>
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<td>87</td>
<td>279</td>
<td>4,855</td>
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<tr>
<td>CSHL</td>
<td>306</td>
<td>337</td>
<td>417</td>
<td>2,887</td>
<td>223</td>
<td>486</td>
<td>5,146</td>
</tr>
<tr>
<td>IoSUGA</td>
<td>919</td>
<td>330</td>
<td>1,665</td>
<td>2,933</td>
<td>356</td>
<td>109</td>
<td>6,310</td>
</tr>
<tr>
<td>nGLC</td>
<td>23</td>
<td>64</td>
<td>2,350</td>
<td>2,237</td>
<td>68</td>
<td>2,532</td>
<td>7,283</td>
</tr>
<tr>
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<td>757</td>
<td>730</td>
<td>905</td>
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<td>216</td>
<td>4,722</td>
<td>6,522</td>
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<tr>
<td>nVelv</td>
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<td>1,473</td>
<td>2,838</td>
<td>306</td>
<td>669</td>
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<tr>
<td>CIuGc</td>
<td>1,205</td>
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<td>1,189</td>
<td>2,026</td>
<td>65</td>
<td>6,113</td>
<td>11,282</td>
</tr>
<tr>
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<td>2,731</td>
<td>2,396</td>
<td>5,908</td>
<td>6,223</td>
<td>1,018</td>
<td>6,711</td>
<td>24,987</td>
</tr>
</tbody>
</table>

Table 6 — Inclusion of annotated features within perfect paths. Each annotation is represented as a set of maximal non-overlapping intervals upon the haplotypes of α 1,2. Each column represents an annotation type, giving the number of bp contained within intervals of the type that are fully contained within perfect paths, as a proportion of all bp in intervals of the type. Annotations from left to right: Full length gene transcripts, exons, untranslated regions (UTRs), non-coding conserved elements and repeats.

<table>
<thead>
<tr>
<th>ID</th>
<th>COG-x shoots (996,462)</th>
<th>CO-o eds (362,627)</th>
<th>CO-u tr (233,835)</th>
<th>CO-n ax+nce (21,292,660)</th>
<th>CO-repeat (14,475,489)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASTR</td>
<td>0.11</td>
<td>0.02</td>
<td>0.82</td>
<td>0.92</td>
<td>0.56</td>
</tr>
<tr>
<td>WTDS-P</td>
<td>0.09</td>
<td>0.06</td>
<td>0.82</td>
<td>0.92</td>
<td>0.59</td>
</tr>
<tr>
<td>EBI</td>
<td>0.08</td>
<td>0.07</td>
<td>0.70</td>
<td>0.99</td>
<td>0.55</td>
</tr>
<tr>
<td>WTDS-S</td>
<td>0.07</td>
<td>0.09</td>
<td>0.75</td>
<td>0.99</td>
<td>0.56</td>
</tr>
<tr>
<td>CRACCS</td>
<td>0.07</td>
<td>0.09</td>
<td>0.72</td>
<td>0.97</td>
<td>0.53</td>
</tr>
<tr>
<td>BCCGSC</td>
<td>0.08</td>
<td>0.04</td>
<td>0.70</td>
<td>0.99</td>
<td>0.59</td>
</tr>
<tr>
<td>DOE/GI</td>
<td>0.05</td>
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<td>0.65</td>
<td>0.99</td>
<td>0.48</td>
</tr>
<tr>
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</tr>
<tr>
<td>CSHL</td>
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<td>0.04</td>
<td>0.80</td>
<td>0.99</td>
<td>0.57</td>
</tr>
<tr>
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<td>0.99</td>
<td>0.42</td>
</tr>
<tr>
<td>IoSUGA</td>
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<td>0.07</td>
<td>0.81</td>
<td>0.99</td>
<td>0.58</td>
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<tr>
<td>UCSF</td>
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<td>0.04</td>
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<td>0.96</td>
<td>0.37</td>
</tr>
<tr>
<td>RHUL</td>
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<td>0.06</td>
<td>0.81</td>
<td>0.99</td>
<td>0.59</td>
</tr>
<tr>
<td>GACWT</td>
<td>0.02</td>
<td>0.07</td>
<td>0.37</td>
<td>0.88</td>
<td>0.38</td>
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<tr>
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<td>0.49</td>
<td>0.80</td>
<td>0.39</td>
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<tr>
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<td>0.84</td>
<td>0.99</td>
<td>0.64</td>
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<tr>
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<td>0.83</td>
<td>0.99</td>
<td>0.64</td>
</tr>
<tr>
<td>nVelv</td>
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<td>0.08</td>
<td>0.89</td>
<td>0.99</td>
<td>0.34</td>
</tr>
<tr>
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<td>0.70</td>
<td>0.99</td>
<td>0.41</td>
</tr>
<tr>
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<td>0.01</td>
<td>0.78</td>
<td>0.99</td>
<td>0.46</td>
</tr>
</tbody>
</table>

Table 7 — Summary of metrics used in the analysis.
<table>
<thead>
<tr>
<th>Description</th>
<th>Unit</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>N50</td>
<td>—</td>
<td>A weighted median of the lengths of items, equal to the length of the longest item ( i ) such that the sum of the lengths of items greater than or equal in length to ( i ) is greater than or equal to half the length of all of the items. With regard to assemblies the items are typically contigs or scaffolds.</td>
</tr>
<tr>
<td>NG50</td>
<td>—</td>
<td>Whereas N50 sets the median in relation to the total length of all items in the set, we define NG50 to be normalised by the average length of the ( \alpha_1 ) and ( \alpha_2 ) haplotypes instead of the total length of all sequences as in N50, it is thus more reliable than N50 for comparison between assemblies.</td>
</tr>
<tr>
<td>CPNG50</td>
<td>bp</td>
<td>Contig path NG50. The weighted median of the lengths of contig paths. Contig paths represent maximal subsequences of contigs that are entirely consistent with ( \alpha_{1,2} ).</td>
</tr>
<tr>
<td>SPNG50</td>
<td>bp</td>
<td>Scaffold path NG50. The weighted median of the lengths of scaffold paths. Scaffold paths represent maximal concatenations of contig paths and scaffold breaks that maintain correct order and orientation.</td>
</tr>
<tr>
<td>Structural errors</td>
<td>Counts</td>
<td>An error within a contig or scaffold. Errors include intra and inter chromosomal joins, insertions, deletions, simultaneous insertion and deletions and insertions at the ends of assembled sequences.</td>
</tr>
<tr>
<td>CC50</td>
<td>bp</td>
<td>Correct contiguity 50. The empirically sampled distance between two points in an assembly, where the two points are as likely to be correctly aligned as not.</td>
</tr>
<tr>
<td>Substitution errors</td>
<td>Counts per correct bits</td>
<td>Number of substitution errors per correct bit. Substitution errors are columns in the alignment where the ( \alpha_1 ) and ( \alpha_2 ) haplotypes contain either the same base (homozygous) or different bases (heterozygous) and the alignment contains a base (or IUPAC symbol) different from either ( \alpha_1 ) or ( \alpha_2 ). The metric uses a bit score to allow for IUPAC symbols.</td>
</tr>
<tr>
<td>Copy number errors</td>
<td>Proportions</td>
<td>For a given haplotype column in the MSA the copy number of the simulated genome can be described as an interval ([\text{min}, \text{max}]). Assemblies with copy number outside of this interval are classified either as an excess, for being above the interval, or a deficiency, for being below the interval.</td>
</tr>
<tr>
<td>Coverage</td>
<td>Percent</td>
<td>The coverage is the percent of columns in the MSA of the target (the whole genome, regions of a specific annotation type, etc) that contain positions of the assembly.</td>
</tr>
<tr>
<td>Genic correctness</td>
<td>Percent</td>
<td>The genic correctness is the percentage of bps in spliced transcripts from the haplotype sequences that align to the assembly with 95% coverage using WU-BLAST.</td>
</tr>
</tbody>
</table>