Assessment of human diploid genome assembly with 10x Linked-Reads data

Abstract:
Background: Producing cost-effective haplotype-resolved personal genomes remains challenging. 10x Linked-Read sequencing, with its high base quality and long-range information, has been demonstrated to facilitate de novo assembly of human genomes and variant detection. In this study, we investigate in depth how the parameter space of 10x library preparation and sequencing affects assembly quality, on the basis of both simulated and real libraries.

Findings: We prepared and sequenced eight 10x libraries with a diverse set of parameters from standard cell lines NA12878 and NA24385 and performed whole genome assembly on the data. We also developed the simulator LRTK-SIM to follow the workflow of 10x data generation and produce realistic simulated Linked-Read data sets. We found that assembly quality could be improved by increasing the total sequencing coverage (C) and keeping physical coverage of DNA fragments (CF) or read coverage per fragment (CR) within broad ranges. The optimal physical coverage was between 332X and 823X and assembly quality worsened if it increased to greater than 1,000X for a given C. Long DNA fragments could significantly extend phase blocks, but decreased contig contiguity. The optimal length-weighted fragment length (Wμ_FL) was around 50 – 150kb. When broadly optimal parameters were used for library preparation and sequencing, ca. 80% of the genome was assembled in a diploid state.

Conclusion: The Linked-Read libraries we generated and the parameter space we identified provide theoretical considerations and practical guidelines for personal genome assemblies based on 10x Linked-Read sequencing.

Keywords: 10x Linked-Read sequencing, de novo assembly, diploid human genome, library preparation
We thank the reviewer for these positive comments and address each point below.

That said, I think there are some analyses missing that should be included:

1. I think you should variant call off of the de novo assemblies to see if there are any differences you are missing because you’re only looking at things at a very high structural level.

We have now called SNVs and SVs from our de novo assemblies and from other methods. Please find our results in the responses to points 2-4 of reviewer2.

2. How is phasing affected? I don't see any data on that other than total diploid regions. You should include the changes to the phase block N50. It's mentioned in the abstract, but I don't see it anywhere else.

We have showed the trend of phased block N50 in different linked-read sets in Figure S14, now we also provided the values of phase block N50s in Table S6.

3. Besides NA50 you should include assembly errors such as breakpoints, translocations, inversions, relocations, etc.
   You have a nice dataset here, you should try to get more out of it.

Thank you for the suggestions. We have re-run QUAST and generated several detailed statistics which are now shown in Table S4. These results are consistent with the contig N50s reported in Figure 3.

Minor comments:

58-66, Probably should add this reference for PacBio CCS sequencing, contig N50 is 15 mb, https://www.biorxiv.org/content/10.1101/519025v2
   We have added this reference

65-66, I'd argue that this statement is a bit strong, cost is lowering, and throughput is increasing for these systems
   This is now lines 70-72. We have rephrased the sentence and now write: “However, long-fragment sequencing suffers from extremely high cost (in the case of PacBio CCS), or low base quality (in the case of single-pass reads of either technology), hampering its usefulness for personal genome assembly.”

68 Not a complete sentence
   We fixed this

Ref 27 isn't our stLFR paper, the doi for that is 10.1101/gr.245126.118, and it is commercially available now in some parts of the world
   We have added the new reference and deleted the confusing words in this sentence.

Reviewer #2: Zhang and co-authors present a parameter study for 10x linked-read sequencing experiments with the objective of evaluating the influence of experimentally controllable parameters on the final diploid assembly quality. The authors perform basic performance evaluation in terms of common metrics such as N50 values and provide technical recommendations for designing linked-read sequencing experiments. Additionally, Zhang et al. implemented a software tool for simulating linked-read sequencing data, which they use for parameter assessment given the known (simulated) truth.

While such studies that provide guidance to users of a sequencing technology are very valuable in principle, I have a number of concerns that should be addressed:

1. There is a closely related article by Luo et al. (2017, DOI: 10.1016/j.csbj.2017.10.002) that has been missed. The authors should clarify what the added value of their study is beyond the work by Luo et al. This comment applies to both aspects: guidance to users in terms of 10x sequencing experiments and the utility/features of their data simulation tool (note that Luo et al. also provide a simulator).
We appreciate and cite the work by Luo et al. However, our study provides (1) a more flexible simulation tool and (2) an extensive set of new sequence data.

Regarding (1)
A. We explicitly allow users to input CF, CR, $W_{\mu, FL}$ and $\mu, FL$, which have strong connections with library preparation and Illumina sequencing. For example, CF is driven by input DNA amount and $\mu, FL$ by DNA preparation and potential size selection. LRSIM only lets the user set the total number of reads.
B. The usability of LRTK-SIM is better than LRSIM. LRSIM requires many third party packages and software to be installed first, such as Inline::C perl library, DWGSIM etc. It is not convenient for the users with insufficient computer experience. LRTK-SIM was written in Python and no third-party software was required. It can be installed and gotten started easily. LRTK-SIM can parallel simulate multiple libraries with a variety of parameters simultaneously. The users can compare the performance of different parameters in one run.

Regarding (2)
Luo et al. compared the influence of different parameters by simulation only, which does not always reflect the situation in real sequencing. In our study, we prepared six real libraries with different parameters and could validate our observations from simulation data.

2. The focus of this manuscript is on guiding researchers who are after a cost-effective characterization of individual human genomes. In my view, Zhang et al. should go the full distance and additionally compare to standard Illumina sequencing followed by mapping and variant calling as a baseline. The assembly metrics employed are not so very informative when it comes to the question of which variation (relative to the reference genome) is been missed/captured in standard approaches.

While human assembly is the focus, we believe that much of the interest in our work will come mainly from researchers who are interested in assembling novel genomes. We use human as an assembly model because assembly quality can be gauged by comparison to the reference sequence. Nonetheless ...

Beyond comparing to standard Illumina sequencing, including a detailed comparison to reference-based processing of 10x data (e.g. using LongRanger) would be interesting. In this way, this study would by much more helpful for planning sequencing studies.

... in response to this comment, we now systematically investigate SNV and SV calls from our assemblies. We compare with standard Illumina data and reference-based processing of our 10x data. The standard Illumina data were downloaded from Genome In A Bottle and analyzed with SVABA to generate SV calls, and with BWA and FreeBayes to generate SNV calls. Long ranger was used to generate SNVs and SVs (only deletions) for 10x reference-based analysis. We noted that R9 failed to be analyzed by Long Ranger due to its extremely large CF. We compared SNV and SV calls among the different approaches using vcfeval (https://github.com/RealTimeGenomics/rtg-tools) and truvari (https://github.com/spiralgenetics/truvari), respectively.

For SNVs, we compared the calls from three strategies to the gold standard of NA12878 (ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/NA12878_HG001/latest/GRCh38/) and NA24385 (ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/AshkenazimTrio/HG002_NA24385_son/latest/GRCh38/).

We found that SNVs from reference-based processing of Illumina and 10x data were comparable, and both of them were better than assembly-based SNV calls. For SVs, our assemblies generated many calls that were missed by the reference-based strategy.

We now provide several additional supplementary tables (Table S7-S12) to present these results.
3. The main reason (in my view) for pursuing de novo assembly of human genomes is to access structural variation that is missed otherwise. An evaluation on how much structural variation is (accurately) captured would be of interest to many readers. This is actually something that the authors point out in the Discussion themselves: "Arguably, the metric that matters most in the context of a personal genome is the discovery of variation that lower-cost approaches do not enable."

As implied by the quote, we agree with the reviewer's comment. Consequently, we now compare three linked-read sets from HG002 with the Tier 1 SV benchmark from Genome in a Bottle by using truvari (https://github.com/spiralgenetics/truvari). The results are summarized in Table S13.

4. PacBio CCS reads are available for HG002 (see Wenger et al., http://dx.doi.org/10.1101/519025). Mapping those CCS reads back to your diploid assemblies and calling variants provides an easy and powerful opportunity to assess the sequence quality from an independent technology.

These data became available while our manuscript was in review. We note that the PacBio CCS calls on HG002 are generally reasonably accurate but are not guaranteed to be correct in the absence of a gold standard. Therefore, we prefer to compare them in an overlap analysis with our calls, as opposed to implying that they are a gold standard by using the term "validation". We used vapor (https://github.com/mills-lab/vapor) to validate our SV calls based on PacBio CCS reads from HG002 and include Table S14 to show the validation rates.

Beyond this, your evaluation could be improved by also adding an assembly evaluation perspective that is more biologically motivated, e.g., number of recovered genes/disrupted genes or similar (this should be supported by Quast-LG/BUSCO).

We have added this analysis in Table S4.

Minor comments

- line 51: pedigree based phasing is quite powerful even for trios (where it is able to phase all variants that are homozygous in at least one individual), so I disagree to the statement that this is only feasible in large pedigrees.
We fixed this and removed confusing words.

- lines 60ff: it is unclear which study your are referring to here, please add the citation at the end of the sentence (N50 31.1Mb)
We included a new reference here.

- line 68: broken sentence; also, putting the citation at the end of the sentence increases readability
We fixed this issue.

- lines 71/72: again, unclear which study you are referring to ("Long Fragment Read")
We included a new reference here.

- lines 125ff: is there a specific reason why five and three? (And not, e.g., five and five?) Also, the meaning of L, M, and H in the subscript of L should be explained
Because we generated two additional libraries (L_1L and L_1M for NA12878) to evaluate the effects of CF and CR in assembly, and we believe the trend should be consistent in the two samples. L, M and H represent low, medium and high CF in the experiments. We have clarified this in the manuscript.

- line 129: percent of what?
The percent of GEM in 10x Chromium system.

- line 151: please be more specific about which version of hg38 was used (detail once if identical hg38 was used throughout the rest of the paper [lines 165, 171, 195 and so on...])
The reference was downloaded from 10x website with the version of GRCh38.
Reference 2.1.0.

- line 172: please provide an exact reference for the high confidence regions that you used (e.g., file URL)
  We have added the URL in the manuscript.

- line 208: "in in"
  We fixed this.

- line 208: this sentence is talking about real data, so the reference to Fig 2C and 2D does not match.
  We clarified this in the manuscript.

- line 209: "...but not dramatically... [...] ...appreciably" - this is subjective language, please rephrase and be more fact-oriented (for instance by including the numbers you refer to in parentheses).
  We included the numbers and rephrased the sentence to be more fact-oriented.

- line 250: "_Alignment"?
  We fixed this.

- line 251: what is the denominator for these 91% all bases that are not Ns in the reference genome? (Note that for this analysis, the version of hg38 matters, see comment above).
  "N"s do not contribute to the denominator.

- The authors mention stLFR in line 278. There's a new preprint that's worth citing/discussing: http://dx.doi.org/10.1101/324392
  We have cited their latest version.

- line 296: "extremely long" please say what extremely long means here
  We defined “extremely long” as the DNA fragments longer than 200kb.

- line 570: please be more specific what you mean by "in-house programs", and where the respective sources are available (is that the "Evaluate_diploid_assembly" github?)
  All the source codes for assembly evaluation are available in https://github.com/zhanglu295/Evaluate_diploid_assembly. We added this information in the sentence.

- please add a - preferably open source - license file to your github repositories
  We added the license files in the GitHub.

- "sample prep" is jargon and should be replaced by "sample preparation" (eg. line 41, but also elsewhere)
  We have updated all the “sample prep” to “sample preparation” in the manuscripts.

| Additional Information: |
|-------------------------|
| **Question**            | **Response** |
| Are you submitting this manuscript to a special series or article collection? | No |

**Experimental design and statistics**

Full details of the experimental design and statistical methods used should be given in the Methods section, as detailed in our Minimum Standards Reporting Checklist. Information essential to interpreting the data presented should be made available in the figure legends.
| Have you included all the information requested in your manuscript? | Yes |
|---|---|
| **Resources** | |
| A description of all resources used, including antibodies, cell lines, animals and software tools, with enough information to allow them to be uniquely identified, should be included in the Methods section. Authors are strongly encouraged to cite [Research Resource Identifiers](#) (RRIDs) for antibodies, model organisms and tools, where possible. | Yes |
| Have you included the information requested as detailed in our [Minimum Standards Reporting Checklist](#)? | |
| **Availability of data and materials** | |
| All datasets and code on which the conclusions of the paper rely must be either included in your submission or deposited in [publicly available repositories](#) (where available and ethically appropriate), referencing such data using a unique identifier in the references and in the “Availability of Data and Materials” section of your manuscript. | Yes |
| Have you have met the above requirement as detailed in our [Minimum Standards Reporting Checklist](#)? | |
Assessment of human diploid genome assembly with 10x Linked-Reads data

Lu Zhang¹,²,³, *, Xin Zhou³, *, Ziming Weng², Arend Sidow²,⁴, †

¹Department of Computer Science, Hong Kong Baptist University
²Department of Pathology, Stanford University
³Department of Computer Science, Stanford University
⁴Department of Genetics, Stanford University

*These authors contributed equally to this work. †Correspondence and requests for materials should be addressed to Arend Sidow (email: arend@stanford.edu)
Abstract

Background: Producing cost-effective haplotype-resolved personal genomes remains challenging. 10x Linked-Read sequencing, with its high base quality and long-range information, has been demonstrated to facilitate de novo assembly of human genomes and variant detection. In this study, we investigate in depth how the parameter space of 10x library preparation and sequencing affects assembly quality, on the basis of both simulated and real libraries.

Findings: We prepared and sequenced eight 10x libraries with a diverse set of parameters from standard cell lines NA12878 and NA24385 and performed whole genome assembly on the data. We also developed the simulator LRTK-SIM to follow the workflow of 10x data generation and produce realistic simulated Linked-Read data sets. We found that assembly quality could be improved by increasing the total sequencing coverage ($C$) and keeping physical coverage of DNA fragments ($C_p$) or read coverage per fragment ($C_r$) within broad ranges. The optimal physical coverage was between 332X and 823X and assembly quality worsened if it increased to greater than 1,000X for a given $C$. Long DNA fragments could significantly extend phase blocks, but decreased contig contiguity. The optimal length-weighted fragment length ($W_{\mu FL}$) was around 50 – 150kb. When broadly optimal parameters were used for library preparation and sequencing, ca. 80% of the genome was assembled in a diploid state.

Conclusion: The Linked-Read libraries we generated and the parameter space we identified provide theoretical considerations and practical guidelines for personal genome assemblies based on 10x Linked-Read sequencing.

Keywords: 10x Linked-Read sequencing, de novo assembly, diploid human genome, library preparation
Data description

Introduction

The human genome holds the key for understanding the genetic basis of human evolution, hereditary illnesses and many phenotypes. Whole-genome reconstruction and variant discovery, accomplished by analysis of data from whole-genome sequencing experiments, are foundational for the study of human genomic variation and analysis of genotype-phenotype relationships. Over the past decades, cost-effective whole-genome sequencing has been revolutionized by short-fragment approaches, the most widespread of which have been the consistently improving generations of the original Solexa technology [1, 2], now referred to as Illumina sequencing. Illumina’s strengths and weaknesses are inherent in the sample preparation and sequencing chemistry. Illumina generates short paired reads (2x150 base pairs for the highest-throughput platforms) from short fragments (usually 400-500 base pairs) [3]. Because many clonally amplified molecules generate a robust signal during the sequencing reaction, Illumina’s average per-base error rates are very low.

The lack of long-range contiguity between end-sequenced short fragments limits their application for reconstructing personal genomes. Long-range contiguity is important for phasing variants and dealing with genomic complex regions. For haplotyping, variants can be phased by population-based methods [4, 5] or family-based recombination inference [6, 7]. However, such approaches are only feasible for common variants in single individuals or when a trio or larger pedigree is sequenced. Furthermore, highly polymorphic regions such as the HLA in which the reference sequence does not adequately capture the diversity segregating in the population are refractory to mapping-based approaches and require de novo assembly to reconstruct [8]. Short-read/short-fragment data are challenged by interspersed repetitive sequences from mobile elements and by segmental duplications, and only support highly fragmented genome reconstruction [9, 10].
In principle, many of these challenges can be overcome by long-read/long-fragment sequencing [11, 12]. Assembly of Pacific Biosciences (PacBio) or Oxford Nanopore (ONT) data can yield impressive contiguity of contigs and scaffolds. In one study [13], scaffold N50 reached 31.1Mb by hierarchically integrating PacBio long reads and BioNano for a hybrid assembly, which also uncovered novel tandem repeats and replicated the structural variants that were newly included in the updated hg38 human reference sequence. Another study [14] produced human genome assemblies with ONT data, in which a contig N50 ~3Mb was achieved, and long contigs covered all class I HLA regions. A recent whole genome assembly of NA24385 [15] with high quality PacBio CCS reads generated contigs with an N50 of 15Mb. However, long-fragment sequencing suffers from extremely high cost (in the case of PacBio CCS), or low base quality (in the case of single-pass reads of either technology), hampering its usefulness for personal genome assembly.

Hierarchical assembly pipelines in which multiple data types are used as another approach for genome assembly [16]. For example, in the reconstruction of an Asian personal genome, fosmid clone pools and Illumina data were merged, but because fosmid libraries are highly labor intensive to generate and sequence, this approach is not generalizable to personal genomes. The "Long Fragment Read" (LFR) approach [17], where a long fragment is sequenced at high depth via single-molecule fragmented amplification, reported promising personal genome assembly and variant phasing by attaching a barcode to the short reads derived from the same long fragment. However, because LFR is implemented in a 384 well plate, many long fragments would be labelled by the same barcodes, making it difficult for binning short-reads, and the great sequencing depth required rendered LFR not cost-effective.

An alternative approach is offered by the 10x Genomics Chromium system, which distributes the DNA preparation into millions of partitions where partition-specific barcode sequences are
attached to short amplification products that are templated off the input fragments. Because of
the limited reaction efficiency in each partition, the sequencing depth for each fragment is too
shallow to reconstruct the original long-fragment, distinguishing this approach from LFR [18].
However, to compensate for the low read coverage of each fragment, each genomic region is
covered by hundreds of DNA fragments, giving overall sequence coverage that is in a range
comparable to standard Illumina short-fragment sequencing while providing very high physical
coverage. Novel computational approaches leveraging the special characteristics of 10x
Genomics data have already generated significant advances in power and accuracy of
haplotyping [19, 20], cancer genome reconstruction [21, 22], metagenomic assemblies [23], and
de novo assembly of human and other genomes [24-26], compared to standard Illumina short-
fragment sequencing. While the uniformity of sequence coverage is not as good as with PCR-
free Illumina libraries, 10x Linked-Read sequencing is a promising technology that combines low
per-base error and good small-variant discovery with long-range information for much improved
SV detection in mapping-based approaches [22, 27], and the possibility of long-range contiguity
in de novo assembly [24, 26, 28].

Practical advantages of the technology include the low DNA input mass requirement (1ng per
library, or approximately 300 haploid human genome equivalents). Real input quantities can vary,
along with other factors, to influence an interconnected array of parameters that are relevant to
genome assembly and reconstruction. The parameters over which the experimenter has influence
are (Figure 1): i). $C_R$: average Coverage of short Reads per fragment; ii). $C_F$: average physical
Coverage of the genome by long DNA Fragments; iii). $N_{F/P}$: Number of Fragments per Partition;
iv). Fragment length distribution, several parameters of which are used, specifically $\mu_{FL}$: Average
Unweighted DNA Fragment Length and $W_{\mu_{FL}}$: Length-Weighted average of DNA Fragment
Length. Note that several parameters depend on each other. For example, a greater amount of
input DNA will increase $N_{F/P}$; shorter fragments increase $N_{F/P}$ at the same DNA input amount
compared to longer fragments; less input DNA will (within practical constraints) increase $C_R$ and decrease $C_F$, and their absolute values are set by how much total sequence coverage is generated because $C_R \times C_F = C$.

Our goal in this study was to experimentally explore the 10x parameter space and evaluate the quality of de novo diploid assembly as a function of the parameter values. For example, we set out to ask whether longer input fragments produce better assemblies, or what the effect of sequencing vs. physical coverage is on contiguity of assembly. In order to constrain the parameter space, we first performed computer simulations with reasonably realistic synthetic data. The simulation results suggested certain parameter combinations that we then approximated in the generation of real, high-depth, sequence data on two human reference genome cell lines, NA12878 and NA24385. These simulated and real data sets were then used to produce de novo assemblies, with an emphasis on the performance of 10x’s Supernova2 [24]. We finally assessed the quality of the assemblies using standard metrics of contiguity and accuracy, facilitated by the existence of a gold standard (in the case of simulations) and comparisons to the reference genome (in the case of real data).

Library preparation, physical parameters and sequencing coverage

We made six DNA preparations that varied in fragment size distribution and amount of input DNA, three each from NA12878 and NA24385. From these, we prepared eight libraries, five from NA12878 and three from NA24385 (Table S1). To generate libraries $L_{1L}$, $L_{1M}$ and $L_{1H}$ (the subscripts $L$, $M$ and $H$ represent low, medium and high $C_F$, respectively), genomic DNA was extracted from ca. 1 million cultured NA12878 cells using the Gentra Puregene Blood Kit following manufacturer's instructions (Qiagen, Cat. No 158467). The GEMs were divided into 3 tubes with 5%, 20%, and 75% to generate libraries $L_{1L}$, $L_{1M}$ and $L_{1H}$, respectively (Figure S1-S3). For the
other libraries, to generate longer DNA fragments ($W_{FL} = 150$kb and longer, **Figure S4-S8**), a modified protocol was applied. Two-hundred thousand NA12878 or NA24385 cells of fresh culture were added to 1mL cold 1x PBS in a 1.5 ml tube and pelleted for 5 minutes at 300g. The cell pellets were completely resuspended in the residual supernatant by vortexing and then lysed by adding 200ul Cell Lysis Solution and 1ul of RNaseA Solution (Qiagen, Cat. No 158467), mixing by gentle inversion, and incubating at 37°C for 15-30 minutes. This cell lysis solution is used immediately as input for the 10x Chromium preparation (ChromiumTM Genome Library & Gel Bead Kit v2, PN-120258; ChromiumTM i7 Multiplex Kit, PN-120262). Fragment size of the input DNA can be controlled by gentle handling during lysis and DNA preparation for Chromium. The amount of input DNA (between 1.25 and 4 ng) was varied to achieve a wide range of physical coverage ($C_F$). The Chromium Controller was operated and the GEM preparation was performed as instructed by the manufacturer. Individual libraries were then constructed by end repairing, A-tailing, adapter ligation and PCR amplification. All libraries were sequenced with three lanes of paired-end 150bp runs on the Illumina HiSeqX to obtain very high coverage ($C=94x-192x$), though the two with the fewest number of gel beads ($L_{LL}$ and $L_{LM}$) exhibited high PCR duplication rates because of the reduced complexity of the libraries (**Table S1**).

**Linked-Reads subsampling**

The high sequencing coverage in the libraries allowed subsampling to facilitate the matching of parameters among the different libraries, for purposes of comparability; these subsampled Linked-Read sets are denoted $R_{id}$ (**Figure 1**). We aligned the 10x Linked-Reads to human reference genome (hg38, GRCh38 Reference 2.1.0 from 10x website) followed by removing PCR duplication by barcode-aware analysis in Long Ranger[21]. Original input DNA fragments were inferred by collecting the read-pairs with the same barcode that were aligned in proximity to each other. A fragment was terminated if the distance between two consecutive reads with the identical
barcode larger than 50kb. Fragments were required to have at least two read pairs with the same barcode and a length of at least 2 kb. Partitions with fewer than three fragments were removed. We subsampled short-reads for each fragment to satisfy the expected $C_R$.

**Generating 10x simulated libraries by LRTK-SIM**

To compare the observations from real data with a known truth set, we developed LRTK-SIM, a simulator that follows the workflow of the 10x Chromium system and generates synthetic Linked-Reads like those produced by an Illumina HiSeqX machine (Supplementary Information and Figure S9). Based on the parameters commonly employed by 10x Genomics Linked-Read sequencing and the characteristics of our libraries, LRTK-SIM generated simulated datasets from the human reference (hg38), explicitly modeling the five key steps in real data generation. Parameters in parentheses are from the standard 10x Genomics protocol: 1. Shearing genomic DNA into long fragments ($W_{FL}$ from 50kb to 100kb); 2. Loading DNA to the 10x Chromium instrument (~1.25ng DNA); 3. Allocating DNA fragments into partitions which are attached the unique barcodes (~10 fragments per partition); 4. Generating short fragments; 5. Generating Illumina paired-end short reads (800M~1200M reads). LRTK-SIM first generated a diploid reference genome as a template by duplicating the human reference genome (hg38) into two haplotypes and inserting SNVs from high-confidence regions in GIAB of NA12878 (ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/NA12878_HG001/latest/GRCh38/HG001_GRCh38_GIAB\_highconf\_CG-III\_FB-III\_GATKHC-Ion-10X-SOLID\_CHROM1-\_X_v.3.3.2_highconf\_nosomaticdel\_noCENorHET7.bed); For low-confidence regions we randomly simulated 1 SNV per 1 kb. The ratio was 2:1 for heterozygous and homozygous SNVs. From this diploid reference genome, LRTK-SIM generated long DNA fragments by randomly shearing each haplotype with multiple copies into pieces whose lengths were sampled from an exponential distribution with mean of $\mu_{FL}$. These fragments were then allocated to pseudo-
partitions, and all the fragments within each partition were assigned the same barcode. The number of fragments for each partition was randomly picked from a Poisson distribution with mean of $N_{F/P}$. Finally, paired-end short reads were generated according to $C_R$ and replaced the first 16bp of the reads from forward strand to the assigned barcodes followed by 7 Ns. More information about implementation can be found in **Supplementary Information**. From that diploid genome, Linked-Read datasets were generated that varied in $C_R$, $C_F$ and $\mu_{FL} (\mu_{FL})$ (**Table S2-S3**). Varying $N_{F/P}$ was only done for chromosome 19 because of the infeasibility of running Supernova2 on whole genome assemblies with large $N_{F/P}$; within practically reasonable values, $N_{F/P}$ does not appear to influence assembly quality (**Figure S10**). In total, we generated 17 simulated Linked-Read datasets to explore the overall parameter space (**Table S2-S3**) and 11 to match the parameters of the abovementioned real libraries (**Figure 1**).

**Human genome diploid assembly and evaluation**

The scaffolds were generated by the “pseudohap2” output of Supernova2, which explicitly generated two haploid scaffolds, simultaneously. Contigs were generated by breaking the scaffolds if at least 10 consecutive ‘N’s appeared, per definition by Supernova2. For the simulations of human chromosome 19, we used the scaffolds from the “megabubbles” output. Contig and scaffold N50 and NA50 were used to evaluate assembly quality. Contigs longer than 500bp were aligned to hg38 by Minimap2[29]. We calculated contig NA50 on the basis of contig misassemblies reported by QUAST-LG [30]. For scaffolds (longer than 1kb), we calculated the NA50 following Assemblathon 1’s procedure [31] (**Supplementary Information**).

**Genomic variant calls from diploid assembly**

We compared single nucleotide variants (SNVs) and structural variants (SVs) from the diploid regions of our assemblies with the ones from standard Illumina data and reference-based
processing of our 10x data. The standard Illumina data were downloaded from Genome in a Bottle [32] and analyzed with SVABA [33] to generate SV calls, and with BWA [34] and FreeBayes [35] to generate SNV calls. Long ranger (https://support.10xgenomics.com/genome-exome/software/pipelines/latest/what-is-long-ranger) was used to generate SNV and SV (only deletions) calls for 10x reference-based analysis. We noted that R9 failed to be analyzed by Long Ranger due to its extremely large C_F. For SNVs, we benchmarked the calls from three strategies using the gold standard of NA12878 (ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/NA12878_HG001/latest/GRCh38/) and NA24385 (ftp://ftp-trace.ncbi.nlm.nih.gov/giab/ftp/release/AshkenazimTrio/HG002_NA24385/son/latest/GRCh38/). For SVs, we compared three linked-read sets (R9, R10, R11) from HG002 with the Tier 1 SV benchmark from Genome in a Bottle [36] and used VaPoR [37] to validate our SV calls based on PacBio CCS reads from NA24385 (Highly-accurate long-read sequencing improves variant detection and assembly of a human genome). We compared SNV and SV calls among the different approaches using vcfeval [38] and truvari [36], respectively.

**Performance of diploid assembly: influence of total coverage** Diploid assembly by Linked-Reads requires sufficient total read coverage \( C = C_R \times C_F \) to generate long contigs and scaffolds. In this experiment, to explore the roles of both physical coverage \( C_F \) and per-fragment read coverage \( C_R \), we first generated eight simulated libraries whose total coverage \( C \) ranged from 16x to 78x: four with \( C_R \) fixed and increasing \( C_F \) and four with fixed \( C_F \), and increasing \( C_R \) (Table S2). Contig and scaffold N50s increased along with increasing either \( C_F \) or \( C_R \) (Figure 2A and 2B). To investigate whether the trend was also present in the real datasets, we analyzed six real libraries (three by varying \( C_F \), and the other three by varying \( C_R \); Figure 1): as \( C \) increased, we varied \( C_F \) and \( C_R \) independently by fixing the other parameter. Contig and scaffold N50s also increased in these simulation (Figure 2C and 2D) and real linked-read sets (Figure 2E and 2F) as a function of total coverage \( C \). Contig lengths did increase a little (621.4kb to 758.1kb for
simulation; 110.7kb to 119.6kb for real data) when $C$ was increased beyond 56X. Accuracy, which we define as the ratio between NA50 (N50 after breaking contigs or scaffolds at assembly errors) and N50 (Figure 2C and 2E), changed 18% for simulation and 7% for real data (587.5kb to 713.3kb for simulation; 97.1kb to 104.5kb for real data). For scaffolds in the real data sets, when $C$ increased from 48X ($R_3$) to 67X ($R_4$), both scaffold N50 and NA50 were significantly improved (N50: 13.4Mb to 30.6Mb; NA50: 6.3Mb to 12.0Mb), but the accuracy dropped slightly from 46.6% to 39.1%, which indicated that scaffold accuracy may be refractory to extremely high $C$ (Figure 2F). These results indicated that assembly length and accuracy were comparable over a broad range of $C_F$ and $C_R$ at constant $C$, which implied that assembly quality was mainly determined by $C$.

**Performance of diploid assembly: influence of fragment length and physical coverage.** To investigate if input weighted fragment length (as measured by $W_{\mu FL}$) influenced assembly quality, we generated four simulated libraries (Table S3) with fixed $C_F$ and $C_R$ and a range of fragment lengths (Figure 3A). Contig length decreased with increasing fragment length, a trend that was also seen in six real libraries (Figure 3B; $C=56X; R_6$ to $R_{11}$ in Figure 1). We then simulated another six libraries with the same parameters as the real ones to explore the effects of physical coverage at constant $C=56x$ (Figure 3C). Contig lengths decreased as a function of increasing physical coverage, a trend that is somewhat less clear in real data possibly due to confounding other parameters such as fragment length (Figure 3D). The two linked-read sets with the worst contig qualities in NA12878 ($R_7$) and NA24385 ($R_{10}$) also showed a significant increase of the number of breakpoints (Table S4).

**Performance of diploid assembly: nature of the source genome.** Assembly errors may occur because of heterozygosity, repetitive sequences, or sequencing error. To illuminate possible
sources of assembly error, we performed simulations by generating 10x-like Linked-Reads as above from human chromosome 19, and then quantified assembly error against these synthetic gold standards. Removal of interspersed repeat sequences from the source genome resulted in better contigs with no loss of accuracy in experiments by varying $C_F$, $C_R$ and $\mu_{FL}$ (Figure 4A, 4C and 4E) and better scaffolds only if $C_R$ was above 1X (Figure 4D). Removal of variation had little effect on contigs and only gave rise to longer scaffolds if $C_R$ was above 0.8X (Figure S11), which is difficult to achieve with real libraries. Finally, a 1% uniform sequencing error had no discernible effect (Figure S12).

**Performance of diploid assembly: fraction of genome in diploid state.** While contiguity is an important parameter for any whole genome assembly, evaluation of diploid assemblies necessitates estimating the fraction of the genome in which the assembly recovered the diploid state. To this end, we divided the contigs generated by Supernova2 into “diploid contigs”, which were extracted from its megabubble structures, and “haploid contigs” from non-megabubble structures. Pairs of scaffolds were extracted as the two haplotypes from megabubble structures if they shared the same start and end nodes in the assembly graph. Diploid contigs were generated by breaking the candidate scaffolds at the sequences with least 10 consecutive ‘N’s and were aligned to human reference genome (hg38) by Minimap2. The genome was split into 500bp windows and diploid regions were defined as the maximum extent of successive windows covered by two contigs, each from one haplotype. Alignment against the human reference genome revealed the overall genome coverages of the six assemblies to be around 91%. For most assemblies, 70%-80% of the genome was covered by two homologous contigs (Table 1), with $R_o$ only reaching 58.9%, probably due to the short fragments of the DNA preparation ($\mu_{FL}=24kb$). We also analyzed another seven assemblies produced by 10x Genomics, all of which had diploid fractions of about 80% as well (Table S5). In the male NA24385, non-pseudoautosomal regions of the X chromosome are hemizygous and should therefore be
recovered as haploid regions. Between 79.9% and 87.6% of these regions were covered by one contig exactly depending on the assembled library. Library construction parameters other than fragment length appeared to have had little impact on the proportion of diploid regions (Tables 1 and Table S5).

Overlapping the diploid regions from the assemblies of the same individual revealed that 50.24% and 67.27% of the genome for NA12878 and NA24385 (Figure S13), respectively, were diploid in all the three assemblies. NA12878 was lower because of the low percentage of diploid regions in assembly *R*₆ (Table 1). The overlaps were significantly greater than expected by chance (NA12878: 33.3%, p-value=0.0049; NA24385: 45.4%, p-value=0.0029. Chi square test). These observations were consistent with heterozygous variants being enriched in certain genomic segments, in which two haplotypes were more easily differentiated by Supernova2. Phase block lengths were mainly determined by total coverage *C* and increased in real data with increasing fragment length (Figure S14, Table S6).

**Performance of diploid assembly: quality of variant calls.** The ultimate goal of human genome assembly is to accurately identify genomic variants. We compared the SNVs and SVs from our assemblies with the calls from referenced-based processing of standard Illumina and 10x data, and benchmarked them using gold standard from Genome in a Bottle and PacBio CCS reads. We found the SNVs from referenced-based processing of standard Illumina and 10x data were comparable and both of them were better than assembly-based calls (Table S7 and S8) For SVs, our assemblies generated many calls that were missed by the reference-based strategy (Table S9-S12) and even by the Tier 1 benchmark of Genome in a Bottle (Table S13), and half of the deletions and a majority of insertions could be validated by PacBio CCS reads (Table S14).
Discussion

In this study, we investigated human diploid assembly using 10x Linked-Read sequencing data on both simulated and real libraries. We developed the simulator LRTK-SIM to examine the likely impact of parameters in diploid assembly and compared results from simulated reads to those from real libraries. We thus determined the impact of key parameters (\(C_R\), \(C_F\), \(N_{F/P}\) and \(\mu_{FL}/\mu_{FL}\)) with respect to assembly continuity and accuracy. Our study provides a general strategy to evaluate assemblies of 10x data and may have implications for the evaluation of other barcode-based sequencing technologies such as CPTv2-seq [39] or stLRF [40] in the future.

10x Practicalities

For standard Illumina sequencing, library complexity is usually sufficient to generate tremendous numbers of reads from unique templates and read coverage can be increased simply by sequencing more. However, the 10x Chromium system performs amplification in each partition, and generally only about 20% to 40% of the original long fragment sequence can be captured as short fragments and eventually as reads, resulting in shallow sequencing coverage per fragment. Sequencing more deeply does not increase the per-fragment coverage much as most of the extra reads are from PCR duplicates. The solution is to sequence multiple 10x libraries constructed from the same DNA preparation and merge them for analysis. This means that \(C_R\) remains in the standard range where PCR duplicates are relatively rare, but \(C_F\) increases proportionally to the number of libraries used. A practical limitation to this approach is that Supernova2 limits the number of barcodes to 4.8 million.

Our results showed that in practice, \(C_F\) should be between 335X and 823X, but no larger than 1000X, given the optimal coverage of \(C=56X\) recommended by 10x and the requirement for sufficient per-fragment read coverage. Surprisingly, we observed that including more extremely
long fragments was detrimental for assembly quality. This is possibly due to the loss of barcode specificity for fragments spanning repetitive sequences. From a computational perspective, too many long fragments are harmful to deconvolving the de bruijn graph, as more complex paths need to be picked out. In our experiments, \( W_{\mu FL} \) between 50kb and 150kb is the best choice to generate reliable assemblies.

### Parameters driving assembly quality

Our results regarding assembly quality, and the 10x parameters that influence it, may be useful for efforts in which de novo assemblies are important for generation of an initial reference sequence. We show that maximization of N50 does not necessarily reflect assembly quality, which we were able to compare to NA50 because there exists a high-quality human reference genome. Contig and scaffold lengths mostly increased with ascending sequencing coverage, and at sufficient overall sequence coverage it did not matter much whether the increasing coverage \( C \) was accomplished by increasing \( C_R \) or \( C_F \). However, both contig and scaffold accuracy decreased with increasing \( C \). We also found, counterintuitively, that contig and scaffold length mostly decreased with increasing fragment length, a phenomenon that may be due to the specific implementation; however, until there is another assembler that can be compared to Supernova2 it will not be possible to reason about this effect. In addition, intrinsic properties of the genome matter greatly, as removal of repeats or lack of variation dramatically improves assembly quality.

Diploid assembly is the appropriate approach for assembly of genomes of diploid organisms that harbor variation. Therefore, an important metric to evaluate diploid assembly is the fraction of the genome that is assembled in a diploid state. The short input fragment length of \( \delta_6 \) resulted in roughly 20% less of the genome in a diploid state (<60% vs <80%) compared to the other libraries of the same individual. This observation suggests that in addition to metrics such as N50,
evaluation of assembly quality should also include the fraction of the genome (or the assembly) that is in a diploid state.

**Cost-benefit analysis**

Overall, we have attempted to give practical guidelines to assembly of 10x data with Supernova2 and evaluate the performance across a wide range of metrics. Arguably, the metric that matters most in the context of a personal genome is the discovery of variation that lower-cost approaches do not enable. We estimate that the cost increase over standard Illumina sequencing is about 2x, given the 10X preparation cost and the higher level of sequence coverage required. There may be many applications for which this combination of excellent single nucleotide variant detection (via barcode-aware read mapping) and precise structural variant discovery (via assembly), achieved by the same data set, is worth the price.

**Comparison with hybrid assemblies**

Hybrid assembly strategies have been applied successfully to produce human genome assembly of long contiguity [13, 14, 41]. In these studies, long contigs are first produced by single-molecule long-reads, such as PacBio (NG50=1.1Mb; [13]) or Nanopore (NG50=3.21Mb; [14]) comparing favorably to our best results for Linked-Reads assemblies (NG50=236kb). Scaffolding is then performed with complementary technologies such as BioNano to capture chromosomal level long-range information. It promoted the scaffold N50 of PacBio to 31.1Mb [13] and Illumina mate-pair sequencing with 10x data to 33.5Mb [25]. Using SuperNova2, the scaffold N50 from our studies reached ~27.86Mb ($R_6$) on the basis of 10x data alone, suggesting that 10x technology gives broadly comparable results at a fraction of the price of long-read-based hybrid assemblies.
Availability of supporting data

The raw sequencing data are deposited in the Sequence Read Archive and the corresponding BioProject accession number is PRJNA527321. Diploid assemblies and the codes for comparison are currently available at http://mendel.stanford.edu-supplementarydata/zhang_SN2_2019 and https://github.com/zhanglu295/Evaluate_diploidAssembly. LRTK-SIM is publicly available at https://github.com/zhanglu295/LRTK-SIM.

Additional files

Table S1. Parameters of libraries prepared for NA12878 and NA24385.  
Table S2. Parameters used to generate linked-read sets for evaluating the impact of $C_F$ and $C_R$ on assemblies.  
Table S3. Parameters used to generate linked-read sets for evaluating the impact of $\mu_{FL}$ and $N_{F,P}$ on assemblies.  
Table S4. Contig misassemblies and recovered transcripts of the six assemblies.  
Table S5. Genomic coverage and fraction of contigs in diploid state generated by Supernova2 for the seven libraries prepared by 10x Genomics. Non-PAR: non-pseudoautosomal regions of X chromosome. WFU, YOR, YORM, PR are female; HGP, ASH and CHI are male.  
Table S6. Phase block N50s of the six assemblies.  
Table S7. Comparison SNV calls from standard Illumina data, 10x reference-based calls, and assembly-based calls for NA12878. All calls were compared to the Genome in a Bottle benchmark.  
Table S8. Comparison SNV calls from standard Illumina data, 10x reference-based calls, and assembly-based calls for NA24385. All calls were compared to the Genome in a Bottle benchmark.  
Table S9. Comparison of SV calls from standard Illumina data and 10x assembly-based calls for NA12878.  
Table S10. Comparison of SV calls from standard Illumina data and 10x assembly-based calls for NA24385.  
Table S11. Comparison of SV calls from 10x reference-based and assembly-based calls for NA12878.  
Table S12. Comparison of SV calls from 10x reference-based and assembly-based calls for NA24385.
Table S13. Comparison of SV calls from our de novo assemblies with the Tier 1 SV benchmark from Genome in a Bottle.

Table S14. Proportion of assembly-based SV calls supported by PacBio CCS reads.

Figure S1. Basic statistics for $L_{1U}$. The distributions of A. the number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S2. Basic statistics for $L_{1M}$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S3. Basic statistics for $L_{1H}$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S4. Basic statistics for $L_2$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S5. Basic statistics for $L_3$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S6. Basic statistics for $L_4$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.
Figure S7. Basic statistics for $L_5$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S8. Basic statistics for $L_6$. The distributions of A. number of fragments per partition; B. sequencing depth per fragment; C. probability density function of unweighted fragment lengths; D. cumulative density function of unweighted fragment lengths; E. reversed cumulative density function of unweighted fragment lengths; F. reversed cumulative density function of weighted fragment lengths.

Figure S9. The workflow of LRTK-SIM to simulate linked-reads

Figure S10. The effect of $N_{F/P}$ on human diploid assembly of chromosome 19 by Supernova2, where $C$ ($C=60X$; $C_F=300X$ and $C_R=0.2X$) and $\mu_{FL}$ ($\mu_{FL}=37kb$) are fixed.

Figure S11. Comparison of assembly qualities from 10x data with and without single nucleotide variants by changing $C_F$, $C_R$ and $\mu_{FL}$. $C_R$ was fixed to 0.2X in A and B; $C_F$ was fixed to 300X in C and D; $C_R$ was fixed 0.2X and $C_F$ was fixed 300X in E and F.

Figure S12. Comparison of assembly qualities from 10x data with (1% uniform) and without sequencing error by changing $C_F$, $C_R$ and $\mu_{FL}$. $C_R$ was fixed to 0.2X in A and B; $C_F$ was fixed to 300X in C and D; $C_R$ was fixed 0.2X and $C_F$ was fixed 300X in E and F.

Figure S13. Overlaps of diploid regions for the three libraries from the same sample. Diploid regions for NA12878 (A) and NA24385 (B). The percentages denote the proportion of genome is diploid.

Figure S14. Phase block N50s as a function of different parameter combinations. A. simulated linked-reads with predefined parameters (Table S5) by changing $C_F$ and $C_R$; B. simulated linked-reads with matched parameters of real linked-read sets (Table S2) by changing $C_F$ and $C_R$; C. real linked-read sets (Table S2) by changing $C_F$ and $C_R$; D. simulated linked-read sets (Table S3) with different $\mu_{FL}$; E. simulated linked-read sets with matched parameters (Table S3) with real linked-read sets as $C=56X$; F. real linked-read sets with $C=56X$ (Table S3).
**Competing interest**

Arend Sidow is a consultant and shareholder of DNAnexus, Inc.

**Author Contributions**

AS conceived the study. LZ and XZ wrote LRTK-SIM and performed the analyses. ZMW prepared the genomic DNA and 10x libraries. LZ, XZ, ZMW and AS analyzed the results and wrote the paper. All authors read and approved the final manuscript.

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| Linked-reads set | Overall (%) | Diploid regions (%) | Haploid regions (%) | Non-PAR (%) | Total contig length (contig>500bp) | Length of contigs from megabubble (contig>500bp) | Percentage (%) |
|------------------|-------------|---------------------|---------------------|-------------|----------------------------------|-----------------------------------------------|----------------|
| $R_6$            | 91.9        | 58.9                | 27.7                | -           | 5,632,483,053                    | 3,758,345,846                                 | 66.73          |
| $R_7$            | 91.1        | 73.3                | 11.3                | -           | 5,613,140,437                    | 4,668,186,478                                 | 83.17          |
| $R_8$            | 91.7        | 77.2                | 9.2                 | -           | 5,635,127,471                    | 4,896,821,850                                 | 86.90          |
| $R_9$            | 91.3        | 73.4                | 12.2                | 85.9        | 5,637,615,919                    | 4,438,175,621                                 | 78.72          |
| $R_{10}$         | 91.7        | 79.2                | 5.8                 | 79.9        | 5,749,001,471                    | 4,793,226,150                                 | 83.37          |
| $R_{11}$         | 91.7        | 78.1                | 7.9                 | 87.6        | 5,677,566,094                    | 4,723,083,367                                 | 83.19          |

Table 1. Genomic coverage of contigs generated by Supernova2. Non-PAR: non-pseudoautosomal regions of X chromosome. $R_6$, $R_7$ and $R_8$ are female; $R_9$, $R_{10}$ and $R_{11}$ are male.
Figure 1. The linked-read sets prepared to evaluate the impact of $C_r$, $C_F$, $\mu_{FL}$ and $W_{\mu_{FL}}$ on human diploid assembly.
**Figure 2.** Contig and scaffold lengths (N50 and NA50) as a function of $C_F$ or $C_R$. **A** and **B**: Simulated Linked-Reads with predefined parameters (Table S2); **C** and **D**: Simulated Linked-Reads with matched parameters of real Linked-Read data sets (Figure 1); **E** and **F**: Real linked-read sets (Figure 1).
Figure 3. Contig qualities (N50 and NA50) as a function of fragment length $W_{\mu_{FL}}$ or physical coverage $C_F$, at $C=56X$. A and C, results from simulations; B and D, results from real data.
Figure 4. Comparison of contig and scaffold lengths from 10x data with masked and unmasked repetitive sequences by changing $C_F$, $C_R$ and $\mu_{FL}$. $C_R$ was fixed to 0.2X in A and B; $C_F$ was fixed to 300X in C and D; $C_R$ was fixed to 0.2X and $C_F$ was fixed to 300X in E and F.
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| Linked-reads set | Overall (%) | Diploid regions (%) | Haploid regions (%) | Non-PAR (%) | Total contig length (contig>500bp) | Length of contigs from megabubble (contig>500bp) | Percentage (%) |
|-----------------|-------------|---------------------|---------------------|-------------|-----------------------------------|-----------------------------------------------|----------------|
| \(R_6\)         | 91.9        | 58.9                | 27.7                | -           | 5,632,483,053                     | 3,758,345,846                                  | 66.73          |
| \(R_7\)         | 91.1        | 73.3                | 11.3                | -           | 5,613,140,437                     | 4,668,186,478                                  | 83.17          |
| \(R_8\)         | 91.7        | 77.2                | 9.2                 | -           | 5,635,127,471                     | 4,896,821,850                                  | 86.90          |
| \(R_9\)         | 91.3        | 73.4                | 12.2                | 85.9        | 5,637,615,919                     | 4,438,175,621                                  | 78.72          |
| \(R_{10}\)      | 91.7        | 79.2                | 5.8                 | 79.9        | 5,749,001,471                     | 4,793,226,150                                  | 83.37          |
| \(R_{11}\)      | 91.7        | 78.1                | 7.9                 | 87.6        | 5,677,566,094                     | 4,723,083,367                                  | 83.19          |

**Table 1.** Genomic coverage of contigs generated by Supernova2. Non-PAR: non-pseudoautosomal regions of X chromosome. \(R_6\), \(R_7\) and \(R_8\) are female; \(R_9\), \(R_{10}\) and \(R_{11}\) are male.
Parameter

- $N_{F/P}$ = Number of fragments per partition
- $\mu_{FL}$ = Mean fragment length
- $W\mu_{FL}$ = Weighted mean fragment length
- $C_R$ = Read coverage per fragment
- $C_F$ = Physical (fragment) coverage
- $C$ = total coverage

Typical values

- 10 - 100
- $\mu_{FL}$ = 10-100kb
- $W\mu_{FL}$ = 20-400kb
- $C_R$ = 0.1x - 0.4x
- $C_F$ = 200x - 1000x
- $C = C_R * C_F = 40x - 80x$

| Linked-read set | Sequenced Library | $\mu_{FL}$ (kb) | $W\mu_{FL}$ (kb) | $C_F$ (X) | $C_R$ (X) | $C$ (X) |
|-----------------|-------------------|-----------------|-----------------|--------|--------|--------|
| $R_1$ / $S_1$   | $L_{1L}$          | 21.6            | 38.6/35.7       | 19     | 0.2    | 4      |
| $R_2$ / $S_2$   | $L_{1M}$          | 22.4            | 39.7/37.4       | 117    | 0.2    | 24     |
| $R_3$ / $S_3$   | $L_{1M}$          | 22.4            | 39.7/36.8       | 117    | 0.4    | 48     |
| $R_4$ / $S_4$   | $L_{1H}$          | 24.0            | 41.1/40.7       | 334    | 0.2    | 67     |
| $R_5$ / $S_5$   | $L_{1M}$          | 22.4            | 39.7/36.8       | 117    | 0.6    | 72     |
| $R_6$ / $S_6$   | $L_{1H}$          | 24.0            | 41.1/40.6       | 334    | 0.17   | 56     |
| $R_7$ / $S_7$   | $L_2$             | 79.0            | 304.3/131.8     | 123    | 0.45   | 56     |
| $R_8$ / $S_8$   | $L_3$             | 99.2            | 214.5/168.3     | 958    | 0.058  | 56     |
| $R_9$ / $S_9$   | $L_4$             | 92.1            | 216.9/154.1     | 1504   | 0.036  | 56     |
| $R_{10}$ / $S_{10}$ | $L_5$           | 120.8           | 267.4/203.7     | 208    | 0.27   | 56     |
| $R_{11}$ / $S_{11}$ | $L_6$           | 64.2            | 151.7/107.6     | 803    | 0.07   | 56     |
Figure 3

A. Simulations

B. Real Data

C. Contig N50

D. Contig NA50

WμFL (kb) vs. Length (kb)

WμFL (kb) vs. CF
Supplementary Material

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Dr. Hongling Zhou  
Editor  
GigaScience  

Dear Dr. Zhou,

It is my pleasure to resubmit our revised, significantly improved and extended, manuscript "Assessment of human diploid genome assembly with 10x Linked-Reads data" for your further consideration for publication in GigaScience. We were able to address all of the reviewers' comments, which are addressed point by point in our response, and hope that you will be able to reach a positive decision.

Sincerely,

Arend Sidow, Ph.D.  
Professor of Pathology and of Genetics  
SUMC R353  
Stanford, CA 94305-5324  
arend@stanford.edu  
+1-650-498-7024  
http://www.sidowlab.org  
http://jimb.stanford.edu