SCIENTIFIC DATA

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OPEN Data Descriptor: Extensive and deep sequencing of the Venter/ HuRef genome for developing and benchmarking genome analysis tools

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We produced an extensive collection of deep re-sequencing datasets for the Venter/HuRef genome using the Illumina massively-parallel DNA sequencing platform. The original Venter genome sequence is a veryhigh quality phased assembly based on Sanger sequencing. Therefore, researchers developing novel computational tools for the analysis of human genome sequence variation for the dominant Illumina sequencing technology can test and hone their algorithms by making variant calls from these Venter/HuRef datasets and then immediately confirm the detected variants in the Sanger assembly, freeing them of the need for further experimental validation. This process also applies to implementing and benchmarking existing genome analysis pipelines. We prepared and sequenced 200 bp and 350 bp short-insert wholegenome sequencing libraries (sequenced to 100x and 40x genomic coverages respectively) as well as 2 kb, 5 kb, and 12 kb mate-pair libraries (49x, 122x, and 145x physical coverages respectively). Lastly, we produced a linked-read library (128x physical coverage) from which we also performed haplotype phasing.

Design Type(s)	reference design • protocol optimization design
Measurement Type(s)	whole genome sequencing
Technology Type(s)	DNA sequencing
Factor Type(s)	
Sample Characteristic(s)	Homo sapiens • lymphoblastoid cell line

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Background & Summary

Almost two decades ago the extensive efforts of the Human Genome Project, backed up by work from Celera, resulted in the release of a draft of the first complete sequence of the human genome^{1,2}. This catalyzed a new era of human whole-genome analysis where the now-available human genome sequence has been studied intensely to understand the functions of its parts and their interactions with each other and where a concurrent genome technology revolution has produced ever more powerful platforms to carry out such functional studies³. Since then, increasingly large numbers of human genomes have been sequenced, yielding insights into population-level genetic variation⁴⁻⁶, structural genome variation⁷⁻⁹, and mutational mechanisms¹⁰. Technological advances have progressively improved the information content and reduced the noise profile of sequencing data¹¹. A large variety of methodologies for the routine analysis of sequencing data is now available¹². "Whole-genome sequencing" is now a standing term that refers to the re-sequencing of a given sample of human genomic DNA using, typically, the dominant Illumina DNA sequencing platforms which can quickly produce several hundred million short sequencing reads at affordable costs. These reads are then aligned to the human reference genome and analyzed using various approaches¹²⁻¹⁴, such as mismatch analysis, readdepth analysis, split-read analysis and discordant read-pairs analysis, producing an extensive catalog of sequence variants that are present in the DNA sample in question relative to the human reference sequence. The promise of human genome research is nothing short of a complete transformation of basic life science research, translational research, and eventually the way we diagnose, treat, and find cures for human disease.

It is clear, however, that current standard whole-genome sequence analysis leaves a rather large room for improvement. The standard genome analysis practices of today perform rather poorly in certain contexts, such as in repetitive regions (i.e. in around half the human genome), in the detection and resolution of complex structural variation, or in placing detected variants in their proper haplotypes. Although more advanced and novel computational algorithms that address these limitations are continuously being developed, one essential requirement during this process is that the detected variants are to be experimentally validated in order to establish false-positive rates and to make it possible to further tune and optimize the new algorithms. Experimental validation, especially of complex variants, during the tool development and testing phases is a very laborious and time-consuming process, but it can be circumvented by using a genome for which sufficiently large numbers of variants are already known, i.e. prevalidated. Several studies have been conducted with the goal of extensively characterizing the variants in a small number of human genomes using multiple sequencing technologies^{15,16}. In some human genomes, variants have been carefully and extensively documented, providing a benchmark for other studies^{9,17–20}.

The Venter (HuRef) Genome, however, is especially distinguished for quality among the publiclyavailable human genome sequences as it is the only one for which its complete diploid assembly was generated from high-quality Sanger reads¹⁷ and for which extensive catalogs of SNPs, indels, and structural variation are available^{18,20}. To date, no extensive Illumina sequencing datasets have been available for the Venter/HuRef genome in contrast to other genomes that have been characterized for benchmarking purposes^{15,16}.

To unlock the potential of the Venter/HuRef genome as the outstanding benchmark genome, we have conducted deep whole-genome sequencing (WGS) using a variety of sequencing strategies for the Illumina platform (Table 1). Specifically, we produced short-insert paired-end WGS datasets at a combined sequence coverage of 140x, linked-read data at 42x de-duplicated sequencing coverage (128x physical coverage i.e. the average number of times the genome is spanned by input DNA fragments rather than the average number of times the genome is covered by sequencing reads as in sequencing coverage), and three long-insert (2 kb, 5 kb, and 12 kb) paired-end (i.e. mate-pair) WGS datasets with physical coverages of 49x, 122x, and 145x, respectively (Fig. 1). These datasets are of very high quality (Figs 2–5) and are complemented by the existing Venter/HuRef assembly-quality Sanger reads¹⁷ and long-read sequencing data, which was produced using the Pacific Biosciences platform²¹.

Researchers developing novel computational tools for analyzing whole-genome sequencing data can now test their algorithms by processing the appropriate Venter/HuRef Illumina datasets described here and then turn to the already-available catalogs of sequence variants, or to the original Sanger reads¹⁷, to confirm the characterization of variants detected by their algorithms . Likewise, whenever a laboratory implements a new computational pipeline for human genome analysis, it can now use these Illumina Venter/HuRef datasets to confirm proper implementation and to optimize proper settings for the pipeline.

Methods

Venter/HuRef DNA Sample

The Venter/HuRef DNA sample as obtained as a 50 µg aliquot of LCL-extracted DNA (NS12911) from the Coriell Institute for Medical Research where the iPSC (GM25430) of the same subject is also available (https://catalog.coriell.org/1/HuRef).

Library Type	Library Preparation Method	DNA Input Amount	Number of Read Pairs	PCR Cycles	Read Length	% Duplication	Alignment %
Short-insert (200 bp)	KAPA NGS Hyper Prep Kit	200 nanograms	1579034384	5	$2\times 100 \ bp$	6.66%	98.56%
Short-insert (350 bp)	KAPA NGS Hyper Prep Kit	200 nanograms	632532251	5	$2\times 100 \ bp$	4.10%	99.11%
2 kb Mate Pair	Nextera Mate Pair Sample Preparation Kit	4 micrograms	211613321	10	$2 \times 150 \text{ bp}$	44.22%	92.81%
5 kb Mate Pair	Nextera Mate Pair Sample Preparation Kit	4 micrograms	128097621	5	$2\times150~\text{bp}$	16.39%	94.77%
12kb Mate Pair	Nextera Mate Pair Sample Preparation Kit	8 micrograms	148540181	10	$2 \times 150 \text{ bp}$	68.60%	97.30%
Linked-Read	10X Genomics Chromium Kit	1 nanogram	789239544	8	$2 \times 150 \text{ bp}$	9.85%	89.14%

 Table 1. Summary of library construction and sequencing for short-insert, mate-pair, and linked-read HuRef/Venter WGS libraries.



Figure 1. Schematic diagram of the study. (a) Venter/HuRef genomic DNA was used to generate shortinsert (200 bp and 350 bp), mate-pair (2 kb, 5 kb, and 12 kb), and linked-read libraries. (b) Detailed overview of data generation including bio-sample used, types of Illumina WGS libraries constructed, sequencing instrument platforms, types of sequencing runs, and subsequent analysis of data.

Illumina paired-end WGS

Library Preparation. The library preparation was previously described in detail in Mu *et al.*²⁰. Briefly, 1 μ g of genomic DNA was fragmented using 2 μ L of NEBNext dsDNA fragmentase (New England Biolabs, Ipswich, MA) in 1x fragmentation buffer and 1x BSA. Reaction was kept on ice for 5 minutes



Figure 2. Normalized coverage, GC (%) content windows, base quality at GC (%), and corresponding insert-size histograms for WGS libraries. Normalized coverage (blue squares). Windows at GC% (pink bars). Base Quality at GC% (green line). (a,b) 200 bp short-insert, (c,d) 350 bp short-insert.

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before adding the fragmentase and was incubated at 37 °C for 20 minutes. The reaction was stopped by addition of 5 μ L of 0.5 M EDTA. DNA was purified from the reaction mixture using 0.9x by volume AMPure XP beads (Beckman Coulter, Cat# A63880) and eluted in 50 μ L of 10 mM Tris-Acetate (pH 8.0) buffer. Six independent fragmentation reaction replicates were performed, and the sizes of the DNA were analyzed using Agilent 2100 Bioanalyzer before library preparation.

Library preparation was performed using the KAPA Library Preparation kit (KAPA Biosystems, Wilmington, MA) where 200 ng of fragmented DNA was used as input. Library was constructed according to manufacturer's protocol where the DNA was end-repaired and A-tailed before adapter ligation with Illumina TruSeq Adapter (Index 1). DNA was then purified using 0.8x by volume AMPure XP beads and quantified using the Qubit ds DNA High Sensitivity Assay Kit (Life Technologies, Cat# Q32851). For PCR amplification, 50 ng of DNA was amplified using the KAPA HiFi DNA Polymerase with the following thermocycling conditions: 98 °C/45 s, 5 cycles of (98 °C/15 s, 60 °C/30 s, 72 °C/45 s), 72 °C/1 min, and 4 °C/hold. Primers from the KAPA Library Preparation kit was used for PCR amplification. Afterwards, DNA was purified from the PCR reaction using AMPure XP beads and eluted in 30 μ L of 10 mM Tris-Acetate (pH 8.0) buffer. Six independent experimental replicates were performed, and the purified PCR amplified DNA fragments from each replicate was pooled for size selection and gelpurified from 2% agarose gel. Two size selections were made at 200 bp and 350 bp.

Sequencing. Sequencing of the 200 bp and 350 bp insert-size libraries was described previously in Mu *et al.*²⁰. The libraries were sequenced separately $(2 \times 100 \text{ bp})$ on an Illumina HiSeq 2000 instrument in rapid run mode. For the 200 bp insert-size library, a total of 3,214,626,588 reads generated from 5 sequencing runs was pooled together to obtain 100x genomic coverage. For the 350 bp insert-size library,



Figure 3. Normalized coverage, GC (%) content windows, base quality at GC (%), and corresponding insert-size histograms for mate-pair libraries. Normalized coverage (blue squares). Windows at GC% (pink bars). Base Quality at GC% (green line). (a,b) 2kb-mate-pair, (c,d) 5kb-mate-pair, (e,f) 12kb-mate-pair.





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a total of 1,280,576,580 reads generated from two sequencing runs was pooled together to obtain 40x genomic coverage.

Analysis. Reads were trimmed at the 3' end to a uniform length of 100 bp using FASTX toolkit (http://hannonlab.cshl.edu/fastx_toolkit/; version 0.0.13). The trimmed reads were aligned by BWA-MEM (Li and Durbin 2009; version 0.7.17-r1188) using the hg38 reference with ALT alleles removed (ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/000/001/405/GCA_000001405.15_GRCh38/ seqs_for_alignment_pipelines.ucsc_ids/), and the resulting alignment records were sorted with Samtools (http://www.htslib.org/; version 1.7). Marking of PCR duplicates and calculations of insert-size and coverage information was performed using Picard (http://picard.sourceforge.net; version 2.17.10).



Figure 5. Violin plot of sizes of haplotype blocks constructed using linked-read sequencing (128x physical coverage) for HuRef/Venter Genome for all chromosomes.

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Illumina mate-pair WGS

Library Preparation. Mate Pair libraries at insert sizes 2 kb, 5 kb, and 12 kb were generated from Venter/HuRef DNA using the Nextera Mate Pair Sample Preparation Kit (Illumina, Cat# FC-132-1001) following standard manufacturer's instructions with the exception of the shearing step (see below). The Venter/HuRef DNA sample was first verified as high molecular weight (>15 kb) by running 60 ng, quantified by using the Qubit dsDNA HS Assay Kit (Life Technologies, Cat# Q32851), on 0.8% 1X TAE agarose gel next to the 1 kb Plus DNA Ladder (ThermoFisher Cat# 10787018). Afterwards, for each insert size, 4 µg of the high molecular weight genomic DNA was tagmented with biotinylated junction adapters and fragmented to about 7-8 kb on average in a 400 µL tagmentation reaction containing 12 µL of Tagmentase at 55 °C for 30 min. The tagmented DNA fragments were purified by adding 2X the volume of DNA Binding Buffer with Zymo Genomic DNA Clean & Concentrator Kit (Zymo Research, Cat# D4010) and eluted in 30 uL of Elution Buffer after two washes with the provided Wash Buffer. To fill in the gaps in the DNA adjacent to the junction adapters as a by-product of Tagmentation, single-strand displacement reaction was performed in a 200 µL reaction by adding 132 µL of water, 20 µL of 10x Strand Displacement Buffer, 8 μ L of dNTPs, and 10 μ L of Strand Displacement Polymerase to the 30 μ L elution and at 20 °C for 30 min. DNA purification was then performed in 30 μ L elution with 0.5x volume of AMPure XP Beads (Beckman Coulter, Cat# A63880) and size-selected by using BluePippin (Sage Science). The 0.75% DF 3-10 kb Marker S1 - Improved Recovery and the 0.75% DF 10-18 kb Marker U1 protocols were used for size selection on the BluePippin for insert sizes 5 kb and 12 kb respectively, and 0.75% DF 1-6 kb Marker S1 protocol was used for insert size 2 kb. The "Tight Selection" option was used instead of "Range" for all size selections. The size selected DNA was then circularized overnight (12-16 hours) at 30 °C with Circularization Ligase in a 300 µL reaction.

After overnight circularization, the uncirculated linear DNA was digested by adding $9 \mu L$ of Exonuclease and incubated at 30 °C for 30 minutes and heat inactivated at 70 °C for 30 minutes. Afterwards, 12 μL of Stop Ligation Buffer was added. Circularized DNA was then transferred to T6 (6 × 32 mm) glass tube (Covaris, Part# 520031 and 520042) and sheared *twice* on the Covaris S2 machine (Intensity of 8, Duty Cycle of 20%, Cycles Per Burst of 200, Time of 40 s, Temperature of 2–6 °C). We find that shearing *twice* often creates a tighter final library size distribution which leads to a higher fraction of pass-filter clusters during the Illumina sequencing step.

The mate pair fragments within the sheared DNA fragments contain the biotinylated junction adapter and were selected by binding to Dynabeads M-280 Streptavidin Magnetic Beads (Invitrogen, Part# 112-05D) by adding an equal volume of the Bead Bind Buffer (incubated at 20 °C for 15 minutes on shaking heat block at highest rpm setting). The non-biotinylated molecules in solution were washed away using the Wash Buffer. All downstream reactions were carried out on streptavidin beads with magnetic immobilization and washes with the Wash Buffer between successive reactions (e.g. End Repair, A-Tailing, and Adapter Ligation. The sheared DNA was first End-repaired followed by A-Tailing and TruSeq indexed adapter ligation.

The adapter-ligated DNA was resuspended in 20 μ L of Resuspension Buffer and then PCR amplified in a 50 μ L reaction with 25 μ L of PCR 2X Master Mix and 5 μ L of Primers both provided in the Nextera Mate Pair Sample Preparation Kit (Illumina, Cat# FC-132–1001) to generate the final library. The thermocycling conditions are 98 °C/1 min, 10 cycles of (98 °C/10 s, 60 °C/30 s, 72 °C/30 s), 72 °C/5 min, and 4 °C /hold. The 5 kb mate-pair library was PCR-amplified for 5 cycles instead of 10 cycles. For the 12 kb mate-pair library, 8 μ g of input DNA was used instead of 4 ug. The amplified library (supernatant) was purified using a 0.66x volume of AMPure XP Beads (0.67x vol) and eluted in 20 μ L of Resuspension Buffer. The size distribution of the library was determined by Agilent Technologies 2100 Bioanalyzer (High Sensitivity Assay), and the indexed library concentration was measured by the Qubit dsDNA HS Assay Kit (Life Technologies, Cat# Q32851). **Sequencing**. The Mate-Pair libraries were sequenced on the Illumina NextSeq 500 using the NextSeq 500/550 Mid Output v2 kit (300 cycles) (Illumina, Cat# FC-404-2003) to generate 2 × 151 bp paired-end reads. The libraries were loaded onto the flowcell at a final concentration of 1.8pM and 1% PhiX Control v3 (Illumina, Cat# FC-110-3001). Additional rounds of sequencing also used a final library concentration of 1.8pM and 1% PhiX Control v3.

Analysis. Illumina Nextera Mate Pair junction adapter sequences were first trimmed using NxTrim²² (version 0.4.3) with the "--aggressive --preserve-mp" settings in order to maximize the number of longinsert pairs. Nxtrim outputs four sets of reads, designated "Mate Pair", "Paired-End", "Singleton", and "Unknown." "Mate Pair" reads have junction adapter sequence trimmed off from the 3' end of Read 1 and/or Read 2; "Paired-End" (short-insert) reads have junction adapter sequence trimmed from the 5' end of Read 1 and/or Read 2; "Singleton" reads have junction adapter sequence trimmed from the middle of either Read 1 or Read 2 rendering one of the reads useless. "Unknown" reads have no junction adapter sequences detected. This is most likely because the junction adapter sequence sits in the un-sequenced portion of the template, thus whether reads are "Mate Pair" or "Paired-End" cannot be discerned. Nonetheless, mate-pair reads are present in the "Unknown" fractions as well as paired-end reads. The "Unknown" reads can be used for alignment and analysis if more long-insert information is desired²². Here, the reads designated as "Mate Pair" and "Unknown" were combined, aligned with BWA-MEM²³ against the hg38 reference without ALT alleles (ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/000/001/ 405/GCA_000001405.15_GRCh38/seqs_for_alignment_pipelines.ucsc_ids/), and sorted using samtools (http://www.htslib.org/; version 1.7). Marking of PCR duplicates and calculations of insert-size and coverage information was performed using Picard (http://picard.sourceforge.net; version 2.17.10).

10X Genomics Chromium library for Illumina sequencing

Input genomic DNA preparation. The Venter/HuRef DNA sample (obtained from the Coriell Institute for Medical Research) was first verified as high molecular weight (>15 kb) by running 60 ng, quantified by using the Qubit dsDNA HS Assay Kit (Life Technologies, Cat# Q32851), on 0.8% 1X TAE agarose gel next to the 1 kb Plus DNA Ladder (ThermoFisher Cat# 10787018). Afterwards, 4 μ g of the high molecular weight genomic DNA was loaded on a BluePippin (Sage Science) instrument to select for DNA fragments 30 kb to 80 kb using the "0.75%DF Marker U1 high-pass 30- 40 kb vs3" protocol. The concentration of the selected DNA fragments was then quantified by using the Qubit dsDNA HS Assay Kit (Life Technologies, Cat# Q32851) and diluted to 1 ng/ μ L. The final dilution concentration of 1 ng/ μ L was verified again by performing three technical replicates of Qubit dsDNA HS Assay with 5 μ L of the DNA dilution as input.

Chromium whole-genome linked-read library preparation and sequencing. The linked-read wholegenome library was prepared using the Chromium Genome kit and reagent delivery system (10X Genomics, Pleasanton, CA). The linked-read library was made following standard manufacturer's protocol with 10 cycles of PCR amplification. Briefly 1 ng of DNA (~300 genome equivalents) of sizeselected high molecular DNA was partitioned into ~1.5 million oil droplets in emulsion, tagged with a unique 16 bp barcode within each droplet, and subjected isothermal amplification (30 °C for 3 hours; 65 °C for 30 minutes) by random priming within each droplet.

Amplified (isothermal) DNA was then purified from the droplet emulsion following the manufacturer's protocol using SPRI beads. The purified DNA was then End-Repaired and A-tailed followed by adapter ligation of adapter in the same reaction mixture. DNA was purified from the was the reaction mixture using SPRI beads and eluted in 40 uL. Sample Index PCR amplification (primers and 2X master mix provided in the Chromium Genome kit) was then performed on the eluted DNA in a toal volume of 100 uL with the following thermocycling conditions: 98 °C/45 s, 10 cycles of (98 °C/20 s, 54 °C/ 30 s, 72 °C/20 s), 72 °C/1 min, and 4 °C /hold. Primer index SI-GA-A6 was used. DNA (final linked-read library) was purified from the PCR reaction with SPRI bead size selection following manufacturer's protocol.

Sequencing. The final purified library was quantified by qPCR (KAPA Library Quantification Kit for Illumina platforms, Kapa Biosystems, Wilmington, MA) using the following thermocycling conditions: 95 °C/3 min, 30 cycles of (95 °C/5 s, 67 °C/30 s). The library concentration was calculated in nanomolar (nM) concentration and then diluted to 5 nM. Sequencing (2x151bp, 8 cycles of single indexing) on two lanes of Illumina HiSeq X (flowcell ID: H3MHGALXX, lanes #4 and #5) was performed at Macrogen (Rockville, MD) resulting in a total of 789,239,544 paired reads (Table 1).

Analysis. FASTQ files were generated from raw BCL files using "*mkfastq*" mode in the Long Ranger software (version 2.1.3) from 10X Genomics (Pleasanton, CA). 10X Genomics Chromium library index "SI-GA-A6" was specified in the required sample sheet file for "*mkfastq*". Before alignment, the hg38 genome files were downloaded from ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCA/000/001/405/GCA_000001405.15_GRCh38/seqs_for_alignment_pipelines.ucsc_ids/GCA_000001405.15_GRCh38/seqs_for_alignment_mode in Long Ranger. Sequencing alignment and haplotype phasing were performed using the "*wgs*" mode in Long Ranger, and the options "-*sex* =

NCBI Experiment Accession	Illumina Instrument	File Format	Download Size (GB)	Library
SRX4026204	HiSeq 2000	BAM	63.4	short-insert (200 bp)
SRX1016819*	HiSeq 2000	FASTQ	132.9	short-insert (200 bp)
SRX4026203	HiSeq 2000	BAM	27.9	short-insert (350 bp)
SRX1016818*	HiSeq 2000	FASTQ	51.6	short-insert (200 bp)
SRX3974885	NextSeq 500	BAM	24.7	long-insert (2 kb)
SRX3894584	NextSeq 500	FASTQ	41.9	long-insert (2 kb)
SRX3974886	NextSeq 500	BAM	9.6	long-insert (5 kb)
SRX3894583	NextSeq 500	FASTQ	21.0	long-insert (5 kb)
SRX3974884	NextSeq 500	BAM	13.6	long-insert (12 kb)
SRX3894586	NextSeq 500	FASTQ	25.8	long-insert (12 kb)
SRX3938571	HiSeq X Ten	BAM	79.3	linked-read
SRX3894585	HiSeq X Ten	FASTQ	102.7	linked-read

Table 2. Details of Data Citation 1.

Library	*Deduplicated Coverage	Physical coverage	Mean Read Length After Trimming	Median Paired-End Insert (bp)	Std Dev of Paired-End Insert (bp)	Read 1 % Aligned	Read 2 % Aligned	Read 1 Mismatch Rate	Read 2 Mismatch Rate
Short-insert (200 bp)	92x	99x	r1 100 bp, r2 100bp	216	25	98.86%	98.26%	0.29%	0.39%
Short-insert (350 bp)	36x	66x	r1 100 bp, r2 100bp	367	43	99.40%	98.81%	0.30%	0.46%
2 kb Mate Pair	7x	49x	r1 130 bp, r2 131 bp	1845	293	93.55%	92.07%	2.58%	2.93%
5 kb Mate Pair	6x	122x	r1 118 bp, r2 119 bp	4830	582	95.76%	93.79%	0.90%	1.44%
12kb Mate Pair	3x	145x	r1 126 bp, r2 127 bp	12234	1990	97.72%	96.88%	1.75%	1.94%
Linked-Read	42x	128x	r1 135 bp, r2 151 bp	354	144	91.90%	89.92%	0.94%	1.79%

Table 3. Summary of post sequencing QC, alignment, duplication, coverage and insert-size analysisfor all libraries. *Footnote: Reads with map quality <20 excluded from coverage calculation.</td>

male" and *"--vcmode = freebayes*" were specified. Only *"PASS*" SNPs and Indels 50 bp or smaller were included in the final phased variant vcf.

Data Records

The Venter/HuRef genome sequenced is publicly available through The Coriell Institute for Medical Research (Camden, NJ, USA) both as genomic DNA (catalog ID: NS12911) extracted from lymphoblastoid cell line (LCL) or as retroviral reprogrammed induced pluriplotent stem cell culture (catalog ID: GM25430). As described in the Methods, Venter/HuRef LCL DNA (NCBI SRA biosample accession SAMN03491120) was used for sequencing library preparation in this work.

Illumina short-insert WGS

Approximately 100x sequencing coverage 2x100bp Illumina short-insert (200 bp) WGS data generated from the Illumina HiSeq 2000 is available through NCBI SRA accession SRR7097858 (Data Citation 1) (Table 2). Approximately 40x sequencing coverage 2x100bp Illumina short-insert (350 bp) WGS data generated from the Illumina HiSeq 2000 platform is available through NCBI SRA accession SRR7097859 (Data Citation 1) (Table 2).

Illumina mate-pair WGS

Illumina mate-pair data sequenced (2x150 bp) on the Illumina NextSeq 500 are available through NCBI SRA accessions SRR6951312, SRR6951313, and SRR6951310 for insert sizes 2 kb, 5 kb, and 12 kb respectively (Data Citation 1) (Table 2).

10X Genomics Chromium linked-read Library

10X Genomics Chromium linked-read data sequenced (2x150 bp) on two lanes of the Illumina HiSeq X Ten is available through NCBI SRA accession SRR6951311 (Data Citation 1) (Table 2). The phased variants of the Venter/HuRef genome obtained through the analysis linked reads is available through dbSNP NCBI_ss# 3646580245-3651364986 (For phasing information, request for original submitted vcf file through NCBI dbSNP.) (Data Citation 2).

Mate-Pair Library	Total Reads	Mate Pair	Paired-End	Unknown	Singleton	Mate Pair (%)	Paired-End (%)	Unknown (%)	Singleton (%)
2 kb	276517832	121236869	41396452	110899127	2985384	48.84%	14.97%	40.11%	1.08%
5 kb	169216722	91690504	29429908	45282573	2813737	54.19%	17.39%	26.76%	1.66%
12 kb	183055231	91533027	27584133	62961234	976837	50.00%	15.07%	34.39%	0.53%

 Table 4. Statistics for trimming of Nextera junction adapter sequence using NxTrim for all mate-pair libraries.

Long Ranger Version	2.1.5
Droplet Barcodes Detected	1585712
Mean DNA Per Droplet (bp)	479954
% of Reads with Valid Barcode	94.54%
Mean Barcode Read Quality q-value)	q38
N50 Linked-Reads per Input DNA Molecule	37
Estimated Input DNA Amount (ng)	1.1 ng
Longest Phase Block (bp)	6521210
N50 of Phase Blocks (bp)	924232
Number of Phase Blocks	8882
Input Molecule Length Mean (bp)	32204
Number of Reads	1578479088
Median Insert Size of Aligned Read Pairs	351
Mean Read Depth with PCR duplicates)	61.84
% of non-N bases with 0 coverage	0.06%
% Alignment	89.14%
% PCR Duplication	9.85%
r1 q20 %	94.48%
r1 q30 %	89.18%
r2 q20 %	83.38%
r2 q30 %	70.67%
% phased SNVs	96.70%
heterozygous	2417600
homozygous	1496895
% phased INDELs	93.85%
heterozygous	418735
homozygous	287579

Table 5. Summary of metrics for linked-read sequencing and phasing of the HuRef/Venter genome.

Technical Validation

Illumina short-insert WGS

Sequencing quality of the WGS mate-pair libraries were assessed using FastQC (Supplementary Information). Insert-size, coverage, GC-bias, alignment, and duplication metrics were analyzed using Picard tools (http://broadinstitute.github.io/picard/). These statistics are summarized in Table 1, Table 3 and Fig. 2.

Illumina mate-pair WGS

Sequencing quality of the WGS mate-pair libraries was assessed using FastQC (Supplementary Information). Insert-size, coverage, GC-bias, alignment, and duplication metrics were analyzed using Picard tools (http://broadinstitute.github.io/picard/). These statistics are summarized in Table 1, Table 3 and Fig. 3. Read fractions that were designated by NxTrim²² as "Mate Pair", "Paired-End", "Singletons", and "Unknown" are summarized in Table 4. The "Mate Pair" fraction for all libraries fall within the expected range (~40–60%). Expected for mate-pair libraries, the relatively high rates of PCR duplication (Supplementary Information) result in significant decreases in sequence coverage (3x to 7x) (Table 1, Table 3, Fig. 3). However, the more useful metric for mate-pair sequencing is high physical coverage²⁴.

Here, physical coverage (C_F) is calculated by the equation $C = C_R \times C_F$ where C is the sequencing coverage and C_R is the mean fractional coverage of input DNA fragments. The mean insert sizes for the mate pair libraries are 1.8 kb, 4.8 kb, and 12.2 kb (Table 3, Fig. 3), which results in physical coverage values of 49x, 122x, and 145x respectively. For the 2 kb mate-pair library for an example, C is 7x, and C_R is 0.14 or (130 bp + 131 bp)/1845 bp, thus C_F is 49x (Table 3). The average final library fragment lengths were approximately 800 bp, 800 bp, and 500 bp for the 2 kb, 5 kb and 12 kb mate-pair libraries respectively. The differences in average library fragment lengths most likely contributed to the more extreme tails of the normalized coverage vs GC% for the 2 kb and 5 kb mate-pair libraries (Fig. 3a,c)²⁵.

10X Genomics Chromium Library

Sequencing quality of the linked-read library was assessed using FastQC (Supplementary Information). Input molecule length, coverage, alignment, duplication, droplet barcode, and phasing metrics were analyzed using the Long Ranger software version $2.1.5^{26}$ (Table 1, Table 3, Table 5 and Fig. 4). Overall, 2.4 million and 1.5 million, 0.42 million and 0.29 million heterozygous and homozygous SNVs and indels respectively were called (Table 5). Of which, 96.7% and 93.85% of heterozygous SNVs and Indels respectively were successfully phased in the Venter/HuRef Genome in a total of 8882 haplotype blocks (N50 ~ 0.9 Mbp, longest phase block ~ 6.5 Mbp) (Table 5). Phase blocks for each chromosome are shown in Fig. 5. Similar to mate-pair libraries, the physical coverage of the linked read library is calculated to be 128x from the mean input DNA molecule length of 32 kb.

Usage Notes

The Venter/HuRef genome sequenced in this work is publicly available as both cell line and DNA from Coriell Institute for Medical Research. The mate-pair and linked-read sequencing data used the same DNA sample/extraction as input. It is possible that small differences may exist when compared to the short-insert datasets since the input DNA came from different cell passages and extractions. Researchers are especially encouraged to use the sequencing data in this work in combination with diploid Sanger sequencing data available for the Venter/HuRef genome published in Levy *et al.*¹⁷.

References

- 1. Lander, E. S. et al. Initial sequencing and analysis of the human genome. Nature 409, 860-921 (2001).
- 2. Venter, J. C. et al. The sequence of the human genome. Science 291, 1304-1351 (2001).
- 3. Reuter, J. A., Spacek, D. V. & Snyder, M. P. High-Throughput Sequencing Technologies. Mol. Cell 58, 586-597 (2015).
- 4. The 1000 Genomes Project Consortium *et al.* A map of human genome variation from population-scale sequencing. *Nature* **467**, 1061–1073 (2010).
- 5. The 1000 Genomes Project Consortium *et al.* An integrated map of genetic variation from 1,092 human genomes. *Nature* **491**, 56–65 (2012).
- 6. The 1000 Genomes Project Consortium et al. A global reference for human genetic variation. Nature 526, 68-74 (2015).
- 7. Korbel, J. O. et al. Paired-end mapping reveals extensive structural variation in the human genome. Science 318, 420-426 (2007).
- 8. Sudmant, P. H. et al. An integrated map of structural variation in 2,504 human genomes. Nature 526, 75–81 (2015).
- 9. Chaisson, M. J. P. *et al.* Multi-platform discovery of haplotype-resolved structural variation in human genomes. Preprint at doi: https://doi.org/10.1101/193144 (2017).
- 10. Jónsson, H. *et al.* Parental influence on human germline de novo mutations in 1,548 trios from Iceland. *Nature* 549, 519–522 (2017).
- 11. Kumar, V. et al. Uniform, optimal signal processing of mapped deep-sequencing data. Nat. Biotechnol. 31, 615-622 (2013).
- 12. Pabinger, S. et al. A survey of tools for variant analysis of next-generation genome sequencing data. Brief. Bioinform. 15, 256–278 (2014).
- 13. Alkan, C., Coe, B. P. & Eichler, E. E. Genome structural variation discovery and genotyping. Nat. Rev. Genet. 12, 363-376 (2011).
- 14. DePristo, M. a *et al.* A framework for variation discovery and genotyping using next-generation DNA sequencing data. *Nat Genet* **43**, 491–498 (2011).
- 15. Zook, J. M. et al. Extensive sequencing of seven human genomes to characterize benchmark reference materials. Sci. Data 3, 160025 (2016).
- 16. Eberle, M. A. *et al.* A reference data set of 5.4 million phased human variants validated by genetic inheritance from sequencing a three-generation 17-member pedigree. *Genome Res.* 27, 157–164 (2017).
- 17. Levy, S. et al. The Diploid Genome Sequence of an Individual Human. PLoS Biol. 5, e254 (2007).
- 18. Pang, A. W. et al. Towards a comprehensive structural variation map of an individual human genome. Genome Biol. 11, R52 (2010).
- 19. Parikh, H. et al. svclassify: a method to establish benchmark structural variant calls. BMC Genomics 17, 64 (2016).
- 20. Mu, J. C. *et al.* Leveraging long read sequencing from a single individual to provide a comprehensive resource for benchmarking variant calling methods. *Sci. Rep* **5**, 14493 (2015).
- Lin, M. Comparing de novo assemblies of J. Craig Venter's genome. Figshare doi:https://doi.org/10.6084/m9.figshare.1319564.v1 (2015).
- 22. O'Connell, J. et al. NxTrim: optimized trimming of Illumina mate pair reads. Bioinformatics 31, 2035–2037 (2015).
- 23. Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler transform. *Bioinformatics* 25, 1754–1760 (2009).
- 24. Bishara, A. et al. Read clouds uncover variation in complex regions of the human genome. Genome Res. 25, 1570–1580 (2015).
- 25. Benjamini, Y. & Speed, T. P. Summarizing and correcting the GC content bias in high-throughput sequencing. *Nucleic Acids Res.* **40**, e72–e72 (2012).
- Marks, P. et al. Resolving the Full Spectrum of Human Genome Variation using Linked-Reads. Preprint at doi:https://doi.org/ 10.1101/230946 (2018).

Data Citations

- 1. Zhou, B. & Arthur, J. G. NCBI Sequence Read Archive SRP137779 (2018).
- 2. Zhou, B. NCBI dbSNP ss3646580245-ss3651364986 (2018).

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Author Contributions

B.Z. and R.P. performed the experiments. J.G.A., B.Z., S.S.H., and Y.H. performed the data analysis. B.Z., J.G.A., W.H.W. and A.E.U. conceived of the work. B.Z. & J.G.A. wrote the manuscript.

Additional Information

Supplementary information accompanies this paper at http://www.nature.com/sdata.

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