Improved *de novo* Genome Assembly: Linked-Read Sequencing Combined with Optical Mapping Produce a High Quality Mammalian Genome at Relatively Low Cost

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Abstract:

Current short-read methods have come to dominate genome sequencing because they are cost-effective, rapid, and accurate. However, short reads are most applicable when data can be aligned to a known reference. Two new methods for *de novo* assembly are linked-reads and restriction-site labeled optical maps. We combined commercial applications of these technologies for genome assembly of an endangered mammal, the Hawaiian Monk seal.

We show that the linked-reads produced with 10X Genomics Chromium chemistry and assembled with Supernova v1.1 software produced scaffolds with an N50 of 22.23 Mbp with the longest individual scaffold of 84.06 Mbp. When combined with Bionano Genomics optical maps using Bionano RefAligner, the scaffold N50 increased to 29.65 Mbp for a total of 170 hybrid scaffolds, the longest of which was 84.78 Mbp. These results were 161X and 215X, respectively, improved over DISCOVAR *de novo* assemblies. The quality of the scaffolds was assessed using conserved synteny analysis of both the DNA sequence and predicted seal proteins relative to the genomes of humans and other species. We found large blocks of conserved synteny suggesting that the hybrid scaffolds were high quality. An inversion in one scaffold complementary to human chromosome 6 was found and confirmed by optical maps.

The complementarity of linked-reads and optical maps is likely to make the production of high quality genomes more routine and economical and, by doing so, significantly improve our understanding of comparative genome biology.

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INTRODUCTION:

High quality non-human genomes, especially mammalian genomes, are needed for a variety of reasons including 1) better constraining the limits of allowable nucleotide variation (a useful tool for identifying potentially causative mutations in human disease), 2) the identification of conserved protein and regulatory regions that may explain the morphological or physiological characteristics of different species, 3) establishing the correct relationship of SNPs to genes for association studies, 4) improving our understanding of evolutionary relatedness and mechanisms, and 5) aiding efforts for species conservation and management. The need for fast, economical and high quality genome data for *de novo* genome assembly of rare and endangered species will become especially important as habitat loss, climate change and human impacts accelerate in the 21st century.

Although next generation sequencing has allowed fast and accurate sequencing, to date, the largest hurdle to genome assembly has been the difficulty and cost in obtaining long contiguous sequence scaffolds from short reads. The human genome and those of many model organisms were assembled using methods where large DNA molecules were cloned (e.g., BACs, cosmids, etc.), employed long DNA sequencing technologies (e.g., PacBio), or used augmented short-reads approaches such as mate-pair libraries. Recently, new single-molecule chemistries and analytic methods have been developed to extend these approaches. Here, we have evaluated two technologies for assembling a *de novo* mammalian genome of the Hawaiian monk seal (Neomonarchus schauinslandi), an endangered species endemic to the Hawaiian islands, using techniques that interrogate individual long DNA molecules and have compared these to the short-read based DISCOVAR method (Weisenfeld et al., 2014) which uses overlapping paired-end (PE) Illumina reads. The first method, the 10X Genomics (10XG) Chromium chemistry, incorporates unique molecular indexes (UMI) into single long DNAs and assembles these into gapped scaffolds, based on shared UMIs, using the Supernova software tool (Weisenfeld et al., 2016). The second method used optical maps from Bionano Genomics (Bionano) both as a quality assessment for the Chromium sequences and as a tool to merge the Supernova scaffolds into longer hybrid assemblies.

METHODS:

Sample:

Blood (\sim 10 ml) was collected in EDTA vacutainers during surgery of an adult male seal, shipped on ice packs and processed within two days of collection. The viability of the cells was assessed by trypan blue exclusion. 1 ml of whole blood was stored in LN₂ while 9 ml (1.85x10⁶ cells/ml) were used for lymphocyte separation and subsequent DNA isolation for optical mapping.

DISCOVAR libraries:

1 ug of DNA was used to prepare PCR-free Illumina libraries of ~450 bp mean insert size as described on the Broad Institute website (https://software.broadinstitute.org/software/discovar/blog/?page_id=375). We generated 703 million 250bp paired-end reads on an Illuina HiSeq2500 for the PCR free library, for a read depth of approximately 75x based on estimated genome size.

Analysis was done using DISCOVAR de novo v52488 using default parameters (Weisenfeld et al., 2014).

10X Genomics Chromium:

DNA was isolated using MagAttract (Qiagen) and the molecular weight assayed by pulsed field gel electrophoresis. HMW gDNA concentration was quantitated using a Qubit Fluorometer, diluted to 1.25 ng/ul in TE, and denatured following manufacturers recommendations. Denatured gDNA was added to the reaction master mix and combined with the Chromium bead-attached primer library and emulsification oil on a Chromium Genome Chip. Library preparation was completed following the manufacturer's protocol (Chromium Genome v1, PN-120229). Sequencing-ready libraries were quantified by qPCR (KAPA) and their sizes assayed by Bioanalyzer (Agilent) electrophoresis.

The library was sequenced (151 X 9 X 151) using two HiSeq 2500 Rapid flow cells to generate 975 M reads with a mean read length of 139 bp after trimming. The read 2 Q30 was 87.93% and the weighted mean molecule size was calculated as 92.33 kb. Mean read depth was approximately 61X. The sequence was analyzed using Supernova software (10X Genomics; Weisenfeld et al., 2016) which demultiplexed the Chromium molecular indexes, converted the sequences to fastq files and built a graph-based assembly. The assemblies, which diverge at "megabubbles," consist of two "pseudohaplotypes." The sequence data were originally analyzed using Supernova 1.0 and then repeated using v1.1 which estimates gap sizes rather than introducing an arbitrary value of 100 Ns. As noted above, the Supernova scaffolds were used by the Bionano Hybrid Scaffold tool to create sequence assemblies.

BioNano Genomics:

Optical mapping of large DNA (Xiao et al., 2007) incorporates fluorescent nucleotides at sequence specific sites, visualizes the labeled molecules and aligns these to each other and to a DNA scaffold (Shelton et al., 2015). Lymphocytes were processed following the IrysPrep Kit for human blood with minor modifications. Briefly, PBMCs were spun and resuspended in Cell Suspension Buffer and embedded in 0.6% agarose (plug lysis kit, BioRad). The agarose plugs were treated with Puregene Proteinase K (Qiagen) in lysis buffer (Bionano Genomics) overnight at 50°C and shipped for subsequent processing (S. Brown, KSU). High Molecular Weight (HMW) DNA was recovered by treating the plugs with Gelase (Epicenter), followed by drop dialysis to remove simple carbohydrates. HMW DNA was treated with Nt. BspQI nicking endonuclease (New England Biolabs) and fluorescent nucleotides incorporated by nick translation (IrysPrep Labeling-NLRS protocol, Bionano). Labeled DNA was imaged on the Irys platform (Bionano) and more than 234,000 Mb of image data were collected with a minimum molecule length of 150 kb.

Bionano Genomics Analysis:

Haplotype aware *de novo* assembly was done using the Bionano Genomics RefAligner Assembler (version 5122) software based on the overlap-layoutconsensus paradigm (Xiao et al., 2015; Xiao et al., 2007; Anantharaman et al., 2001; Valouev et al., 2006). To build the overlap-layout-consensus graph, first the single molecule optical maps underwent a pairwise alignment where each molecule was aligned to every other molecule. Pairwise alignments generated using the Bionano Genomics RefAligner were used as input to the layout and consensus assembly stage where a draft assembly was built. We used a P value threshold of 1e-10 for the initial assembly and a minimum molecule length of 150kb. Next we refined the draft assembly using a P value cut off of 1e-11. Refinement of the assembly corrected errors and trimmed and split contigs if errors were found. Our first set of refined contigs were further improved using five iterations of extension and merging during which all contigs were aligned to each other to check for overlap or redundancy; the P value threshold used was 1e-15. Map extension was done by aligning all input molecules to the refined maps. When a set number of molecules extended past the end of a contig, they were combined into a consensus and added to the end of the contig. This increased the size of the contigs. A final refinement step was performed on the maps after the iterative cycles of extension and merging to produce a more accurate and haplotype separated final consensus map. Haplotype separated maps were built by aligning molecules to the genome maps and clustering them into two alleles. When the reported difference between the two alleles was large enough, two haplotype separate maps were generated.

Hybrid scaffolding:

Hybrid scaffolding was performed on the haplotype separated genome maps and the Supernova pseudohaplotype scaffolds. The first step involved in silico digesting the Supernova sequence assembly using the Nt.BspQI recognition motif to generate map coordinates. Next the sequence scaffolds were aligned with the genome maps to flag alignments that were concordant and were fed into the iterative merging stage. The P value threshold used for the initial alignment was 1e-10 while merge was set at 1e-11. After the iterative merge stage the hybrid scaffolds were generated and aligned to the original sequence scaffolds. Finally, we exported back from genome map coordinates to FASTA format along with an AGP file (https://www.ncbi.nlm.nih.gov/assembly/agp/AGP_Specification/) that tracked had been merged.

Scaffold Quality Assessment:

QUAST (Gurevich et al., 2013) was used to generate N50 plots and comparative metrics for the assembly methods. BUSCO (Benchmarking Universal Single Copy Orthologs), a tool for assessing genome completeness (Simão et al., 2015), was run on the hybrid pseudo-reference using 3,023 vertebrate-specific single copy orthologs in genome assembly assessment mode.

Conserved Synteny:

The quality of the Bionano/Supernova hybrid scaffolds was determined in various ways. First, we chose one pseudohaplotype of all 170 unique scaffolds

generated from the Bionano Hybrid Scaffold tool as a pseudo-reference and performed TBLASTN (e.g., Jun et al., 2009) to translate the seal sequences for alignment to both human and dog protein databases. The position of matching proteins from known coding genes in the seal pseudo-reference was manually reviewed in IGV (Thorvaldsdottir et al., 2013). We compared the regions of conserved syntenic genes primarily to the human genome because it is best annotated and, in some instances, to dog, cat and other mammals for which longer scaffolds were available.

DNA alignment:

Conserved syntenic gene order was confirmed using nucmer (Kurtz et al., 2004) to align the seal hybrid pseudo-reference to human (GRCh37/hg19), with a minimum cluster length of 20 and a minimum match length of 50. Plots were generated for all matches greater than 1kb.

Analysis software:

Supplement 1 lists the specific software and commands used for analysis.

RESULTS:

Both 10XG and Bionano software are haplotype aware. For this study we chose Supernova "pseudohaplotype" 1 for analysis. No major differences in length were noted between haplotypes, as might be expected for a species with a population size of about 1400 animals and with reported low heterozygosity (Schultz et al., 2009)

A summary of results is shown in Table 1. The 250bp PE DISCOVAR sequencing

Assembly	BNG_10X_hybrid Supernova 1.0	BNG_10X_hybrid Supernova 1.1	10X Supernova v1.1	DISCOVAR
Total scaffolds	216	170	7,932	437,230
Largest scaffold	70.55 Mbp	84.77 Mbp	84.06 Mbp	1.10 Mbp
N50	20.87 Mbp	29.65 Mbp	22.23 Mbp	137,851
N50 Fold Improvement	105	215	161	1
L50	49	26	36	5100
# N's per 100 kbp	254.65	2,183.45	2,195.29	96.06
Est. total assembly length	2,318 Mbp	2,360 Mbp	2,400 Mbp	2,462 Mbp

Table 1. A comparison of Bionano Irys hybrid scaffolds with 10XG Chromium scaffolds assembled with Supernova v.1.0 (col 2) and v.1.1 (col 3). Col 4 is the 10XG Supernova v1.1 data alone and col 5 is data assembled with DISCOVAR. In addition, 7,696 mostly short scaffolds were unscaffolded with the Irys assembler; their length totaled 40.4 Mbp.

produced over 437,230 reads with a scaffold N50 of 137,851 bp. In contrast, the 10X Chromium sequencing assembled with Supernova v1.0 had an N50 of nearly 14.6 Mbp, an improvement of over 100 fold. When Supernova v1.1 was used the N50 increased to 22.23 Mbp. The Bionano optical maps significantly improved overall scaffold length and decreased the total number of scaffolds from 203 to 170.

Figure 1 shows a QUAST (Gurevich et al., 2013) plot of scaffold size distributions. The QUAST statistics estimated the total assembly length from 2.32-2.46 Gbp, which is similar to that of other carnivores. The total number of N's increased (Table 1) between Supernova v1.0 and v1.1, as expected, due to the improved gap estimation algorithm. The change improved alignment to the optical maps, by more accurately spacing BspQI sites with respect to the sequence scaffolds. Figure 2 shows the corresponding improvement in Bionano confidence scores with Supernova v1.1. Figure 3a shows, in IrysView, an example of a long (~450 kbp) region of high concordance between the optical maps and the Supernova v1.1 scaffold while Fig. 3b shows a region where Supernova may have added larger gaps than appear in the Bionano maps. Figure 4 is an example of where a Bionano map merged two Supernova sequence scaffolds. This example was further studied by conserved synteny analysis.

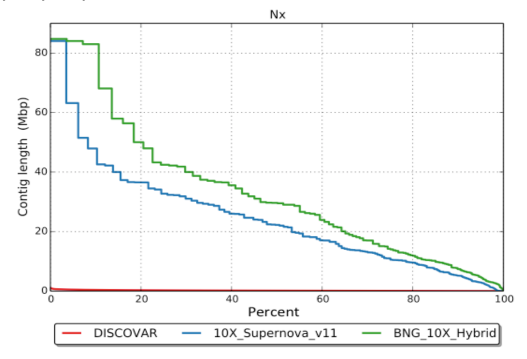


Figure 1. A QUAST (Gurevich et al., 2013) Nx plot of contig lengths as a percent of total scaffolds. The red line represents data assembled by DISCOVAR *de novo* (https://www.broadinstitute.org/software/discovar/blog/; Weisenfeld et al., 2014). Blue shows the improved scaffold lengths using the 10X Genomics Chromium chemistry and Supernova assembler v1.1. The green line is the additional improvement of the Supernova scaffolds when combined with Bionano Genomics optical maps.

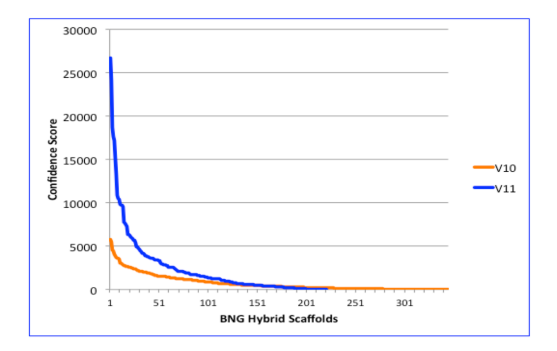


Fig. 2. Improved gap estimation in Supernova v1.1 significantly improved the Bionano confidence scores while reducing the total number of hybrid scaffolds.

In order to identify possible false Bionano joins between Supernova scaffolds, we performed TBLASTN, using a locally installed version of BLAST, to compare the 170 Bionano/10XG hybrid scaffolds against a database of human protein-coding genes. We set the match criteria at low stringency (evalue 1e-05) so that we would maximize the likelihood of identifying homologs but with the understanding that there would be many false matches. We inspected matches by finding two or more contiguous genes that mapped to the same human chromosome and with similar spacing and order. In a few instances of gene families with highly similar paralogs the exactly matching ortholog was not always identified by TBLASTN and in these cases we assigned the specific gene that best fit the conserved synteny relationship.

As an example, Table 2 shows four non-overlapping scaffolds that share conserved synteny with the majority of human chromosome 6. Although we noted inversions in gene order within scaffolds when they were arranged with respect to the chromosome we saw excellent coverage with no large gaps of expected orthologs. When the same analysis was extended to the remainder of the genome (data not shown) we found similar coverage with relatively few gaps indicating that most of the protein-coding genes have been captured in the 170 hybrid scaffolds. The only regions conspicuously absent were the short arms of the acrocentric human chromosome which are largely ribosomal RNA repeats and would not have been identified by TBLASTN analysis. The seal scaffolds orthologous to human chromosome 6 from Fig. 4 are shown in Fig. 5 as a nucmer plot which compares

DNA sequences between seal and human. The plot confirms that the sequence contiguity within the seal scaffolds agrees with the conserved synteny analysis.

We also performed BUSCO analysis of the conserved vertebrate gene set both for the 170 hybrid scaffolds as well as the unscaffolded sequences that did not align to Bionano maps. Of the 170 hybrid scaffolds BUSCO identified 2,696 complete single-copy genes (89.2% of the 3023 expected conserved vertebrate genes) and 16 duplicates (0.5%). These BUSCO assessments are nearly identical to those calculated for humans (Simão et al., 2015). An additional 171 genes were scored as fragmented (5.7%) and 140 were missing (4.6%). We manually searched our TBLASTN predicted protein list and accounted for all but 12 of the "missing." Among the sequences not aligning to optical maps, 8 "complete" and 14 "fragmented" genes were found (fewer than 1% of the total screened).

Table 2. Four seal scaffolds with genes orthologous to human chr 6. The orthologs were ordered according to their conserved syntenic positions on chr 6 and extended the length of the chromosome. The order of the genes highlighted in yellow in scaffold 47 was inverted relative to the human gene order. The inversion was confirmed by Bionano optical maps (Fig. 4).

Scaffold	Position	RefGene	HuChr	Start	End
47	2,841,008	EXOC2	6	485,133	693,111
47	2,953,434	HUS1B	6	655,939	656,963
47	5,432,511	MYLK4	6	2,663,629	2,750,966
47	5,637,247	SERPINB9	6	2,887,266	2,903,280
47	6,850,365	ECI2	6	4,115,689	4,135,597
47	7,644,508	RPP40	6	4,994,732	5,004,063
47	10,025,315	RIOK1	6	7,389,496	7,418,037
47	10,624,264	BLOC1S5	6	8,013,567	8,064,414
47	11,007,567	SLC35B3	6	8,413,068	8,435,483
47	13,360,854	PAK1IP1	6	10,694,695	10,709,782
47	13,481,103	GCM2	6	10,873,223	10,881,941
47	14,288,117	ADTRP	6	11,712,054	11,807,046
47	14,798,305	EDN1	6	12,290,363	12,297,194
47	16,055,443	RANBP9	6	13,621,498	13,711,564
47	16,211,441	MCUR1	6	13,786,549	13,814,994
47	16,500,550	CD83	6	14,117,256	14,136,918
47	17,589,062	JARID2	6	15,246,296	15,522,040
47	18,282,073	MYLIP	6	16,129,125	16,148,248
47	19,523,903	FAM8A1	6	17,600,355	17,611,719
47	20,011,483	KDM1B	6	18,155,329	18,223,853
47	22,093,462	E2F3	6	20,401,906	20,493,715
47	25,658,069	GPLD1	6	24,424,565	24,495,205
47	25,881,304	C6orf62	6	24,704,861	24,720,836
47	26,768,511	SLC17A4	6	25,754,699	25,781,191
47	27,610,706	PRSS16	6	27,247,701	27,256,624
47	28,725,579	GPX6	6	28,503,296	28,528,215
47	29,480,987	GABBR1	6	29,602,228	29,633,135
47	2,917,479	<mark>MOG</mark>	6	29,656,981	29,672,372
47	2,252,098	<mark>HLA-F</mark>	6	29,723,340	29,740,355
47	2,251,182	HLA-A	6	29,942,470	29,945,884
47	2,108,886	<mark>C6orf136</mark>	6	30,647,039	30,653,210

47	1,936,783	DDR1	6	30,870,550	30,889,733
47	1,453,436	DDX3 <mark>9B</mark>	6	31,606,569	31,622,924
47	1,284,137	CLIC1	6	31,730,581	31,737,318
47	1,026,563	TNXB	6	31,998,796	32,043,729
47	1,050,161	CYP21A2	6	32,014,113	32,017,485
47	945,897	PBX2	6	32,133,204	32,138,654
47	932,650	NOTCH4	6	32,194,843	32,224,067
152	38,345	HLA-DRB5	6	32,517,343	32,530,287
152	60,960	HLA-DRB1	6	32,552,990	32,589,848
152	62,928	HLA-DQB1	6	32,659,467	32,668,383
152	62,934	HLA-DPB1	6	33,075,926	33,087,201
152	428,940	COL11A2	6	33,162,681	33,192,499
152	447,468	SLC39A7	6	33,200,445	33,204,439
152	506,118	B3GALT4	6	33,277,132	33,284,832
152	610,185	CUTA	6	33,416,442	33,418,317
152	737,381	BAK1	6	33,572,547	33,580,293
152	793,212	ITPR3	6	33,620,365	33,696,574
152	848,284	UQCC2	6	33,694,293	33,711,727
152	1,166,129	GRM4	6	34,018,645	34,155,622
152	1,502,892	RPS10	6	34,417,454	34,426,125
152	1,584,108	PACSIN1	6	34,466,061	34,535,231
152	1,859,803	UHRF1BP1	6	34,792,015	34,883,138
152	2,066,563	TCP11	6	35,118,071	35,148,610
152	2,343,660	PPARD	6	35,342,558	35,428,191
152	2,369,881	FANCE	6	35,452,361	
	2,369,861 2,417,269				35,467,103
152		TULP1	6	35,497,874	35,512,938
152	2,656,648	ARMC12	6	35,737,032	35,749,079
152	2,744,242	SRPK1	6	35,832,966	35,921,342
152	2,841,367	SLC26A8	6	35,943,514	36,024,868
152	2,994,721	MAPK14	6	36,027,677	36,111,236
152	3,088,997	BRPF3	6	36,196,744	36,232,790
152	3,204,414	C6orf222	6	36,315,757	36,336,885
152	3,348,241	KCTD20	6	36,442,767	36,491,143
152	3,368,929	STK38	6	36,493,892	36,547,470
152	3,577,284	CPNE5	6	36,740,775	36,840,002
152	3,717,442	C6orf89	6	36,871,870	36,928,964
152	3,827,924	FGD2	6	37,005,646	37,029,070
152	4,126,093	RNF8	6	37,353,972	37,394,738
152	4,231,592	CMTR1	6	37,433,131	37,482,844
152	4,821,753	ZFAND3	6	37,819,499	38,154,624
152	5,284,776	BTBD9	6	38,168,451	38,640,148
152	5,363,014	GLO1	6	38,675,925	38,703,141
152	5,418,483	DNAH8	6	38,715,341	39,030,529
152	5,810,207	GLP1R	6	39,048,798	39,087,743
152	6,024,324	KCNK16	6	39,314,698	39,322,968
152	6,047,898	KIF6	6	39,329,990	39,725,405
26	406,011	DAAM2	6	39,792,298	39,904,877
26	2,001,969	FOXP4	6	41,546,426	41,602,384
26	2,206,234	USP49	6	41,789,896	41,895,361
26	2,408,430	TAF8	6	42,050,513	42,087,461
26	2,445,724	C6orf132	6	42,101,118	42,142,619
26	2,508,153	GUCA1B	6	42,184,401	42,194,916
26	2,892,519	UBR2	6	42,564,062	42,693,504
26	3,876,392	VEGFA	6	43,770,184	43,786,487
26	4,372,743	AARS2	6	44,299,654	44,313,326
26	6,042,112	ENPP4	6	46,129,993	46,146,699
26	7,254,492	CD2AP	6	47,477,789	47,627,263
26	9,063,622	MUT	6	49,430,360	49,463,191
26	10,266,365	TFAP2D	6	50,713,828	50,772,988
26	10,345,803	TFAP2B	6	50,818,723	50,847,613

26	11,729,609	MCM3	6	52,264,009	52,284,881
26	13,399,950	MLIP	6	53,929,982	54,266,280
26	14,055,747	FAM83B	6	54,846,771	54,942,022
26	14,562,753	HMGCLL1	6	55,434,369	55,579,214
26	15,552,546	DST	6	56,457,987	56,954,628
26	16,056,305	BEND6	6	56,955,126	57,027,342
26	16,216,175	BAG2	6	57,172,326	57,189,833
26	16,364,502	PRIM2	6	57,314,805	57,646,849
26			6	63,275,951	
	18,733,835	LGSN			63,319,977
26	19,020,962	PHF3	6	63,635,820	63,779,336
26	23,830,984	AGRB3	6	68,635,282	69,389,511
26	25,018,666	FAM135A	6	70,412,941	70,561,174
26	25,658,762	OGFRL1	6	71,288,803	71,308,950
3	84,112,823	RIMS1	6	71,886,703	72,403,143
3	83,186,090	MTO1	6	73,461,578	73,509,236
3	81,949,665	COL12A1	6	75,084,326	75,206,051
3	81,198,935	IMPG1	6	75,921,115	76,072,678
3	78,875,241	IRAK1BP1	6	78,867,472	78,946,440
3	78,731,298	PHIP	6	78,935,867	79,078,236
3	77,882,873	ELOVL4	6	79,914,812	79,947,580
3	76,413,900	FAM46A	6	81,491,439	81,752,774
3	75,114,258	DOPEY1	6	83,067,666	83,171,350
3	75,043,295	PGM3	6	83,161,150	83,193,936
3	74,977,755	ME1	6	83,210,389	83,431,071
3	71,714,453	C6orf163	6	87,344,849	87,365,463
3	71,605,411	SLC35A1	6	87,470,623	87,512,336
3	71,564,591	RARS2	6	87,514,378	87,590,003
3	70,176,222	PM20D2	6	89,146,050	89,165,565
3	69,625,479	MDN1	6	89,642,499	89,819,723
3	66,247,239	EPHA7	6	93,240,020	93,419,547
3	64,349,305	MANEA	6	95,577,543	95,609,457
3	63,458,934	UFL1	6	96,521,595	96,555,276
3	63,160,649	NDUFAF4	6	96,889,313	96,897,881
3	60,825,374	FAXC	6	99,271,169	99,350,062
3	59,606,368	ASCC3	6	100,508,194	100,881,372
3	58,505,231	GRIK2	6	101,398,788	102,070,083
3	55,916,988	HACE1	6	104,728,093	104,859,919
3	55,602,860	POPDC3	6	105,158,280	105,179,995
3	55,407,039	PREP OBSL1	6	105,277,565	105,403,084
3	54,230,441	QRSL1	6	106,629,578	106,668,417
3	53,609,305	PDSS2	6	107,152,557	107,459,683
3	53,229,894	SEC63	6	107,867,756	107,958,278
3	53,324,808	SCML4	6	107,704,104	107,824,317
3	52,662,962	LACE1	6	108,294,874	108,525,784
3	52,242,062	ARMC2	6	108,848,416	108,974,472
3	51,792,347	PPIL6	6	109,390,215	109,441,171
3	51,555,610	FIG4	6	109,691,312	109,825,428
3	50,548,418	AMD1	6	110,874,770	110,895,713
3	49,349,830	LAMA4	6	112,108,760	112,254,939
3	46,185,569	FRK	6	115,931,149	116,060,758
3	45,959,041	NT5DC1	6	116,100,849	116,249,497
3	45,663,548	FAM26F	6	116,461,370	116,463,779
3	44,966,229	ROS1	6	117,288,300	117,425,855
3	43,425,386	MAN1A1	6	119,177,209	119,349,761
3	40,748,527	HSF2	6	122,399,546	122,433,119
3	40,715,709	SERINC1	6	122,443,354	122,471,822
3	40,445,056	SMPDL3A	6	122,789,049	122,809,720
3	37,577,092	TRMT11	6	125,986,430	126,039,276
3	37,313,537	CENPW	6	126,340,174	126,348,875
3	36,442,589	ECHDC1	6	127,288,710	127,343,609

3	34,410,632	LAMA2	6	128,883,141	129,516,569
3	33,701,892	SAMD3	6	130,144,315	130,365,425
3	32,529,868	MED23	6	131,573,966	131,628,229
3	31,526,312	VNN3	6	132,722,787	132,734,765
3	31,491,481	SLC18B1	6	132,769,370	132,798,553
3	30,216,128	SGK1	6	134,169,246	134,318,112
3	29,668,501	ALDH8A1	6	134,917,390	134,950,122
3	29,578,641	HBS1L	6	134,960,378	135,054,898
3	29,426,667	MYB	6	135,181,315	135,219,173
3	28,465,143	BCLAF1	6	136,256,863	136,289,851
3	27,841,899	IL20RA	6	136,999,971	137,045,180
3	27,111,169	TNFAIP3	6	137,867,188	137,883,312
3	26,723,014	ARFGEF3	6	138,161,921	138,344,663
3	25,844,008	CITED2	6	139,371,807	139,374,620
3	23,556,896	NMBR	6	142,058,330	142,088,799
3	22,460,142	AIG1	6	143,060,496	143,340,304
3	22,307,461	PEX3	6	143,450,807	143,490,010
3	21,913,036	PLAGL1	6	143,940,300	144,064,599
3	21,139,050	UTRN	6	144,285,701	144,853,034
3	20,188,841	SHPRH	6	145,864,245	145,964,423
3	19,496,651	ADGB	6	146,598,965	146,815,462
3	18,802,872	SAMD5	6	147,508,927	147,737,547
3	17,136,380	TAB2	6	149,218,641	149,411,613
3	16,889,963	LATS1	6	149,658,153	149,718,256
3	15,547,830	ARMT1	6	151,452,258	151,470,101
3	14,653,210	SYNE1	6	152,121,684	152,637,801
3	14,217,199	MTRF1L	6	152,987,362	153,002,685
3	12,161,992	NOX3	6	155,395,370	155,455,903
3	9,662,930	SERAC1	6	158,109,515	158,168,270
3	8,290,345	TCP1	6	159,778,498	159,789,703
3	8,061,191	IGF2R	6	159,969,099	160,113,507
3	7,775,898	PLG	6	160,702,238	160,753,315
3	5,807,350	PACRG	6	162,727,132	163,315,492
3	3,910,419	C6orf118	6	165,279,664	165,309,607
3	2,290,679	UNC93A	6	167,271,169	167,316,019
3	960,211	THBS2	6	169,215,780	169,254,044
3	39,405	PSMB1	6	170,535,117	170,553,341
3	5,415	TBP	6	170,554,302	170,572,870



Figure 3. a. Example of a long block of high confidence alignment between Bionano optical maps and the Supernova v1.1 sequence. b. Example of where Supernova v1.1 (green arrows) estimated gap sizes that appeared larger than observed optically (blue arrows).

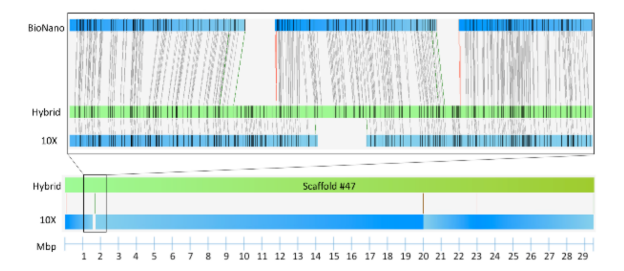


Figure 4. Bionano hybrid scaffold #47 with magnification, above, of the region at approximately 1.7 Mbp. This region is a position of evolutionary breakage or inversion in mammalian genomes. The optical maps confirmed that the DNA scaffolds were correctly joined.

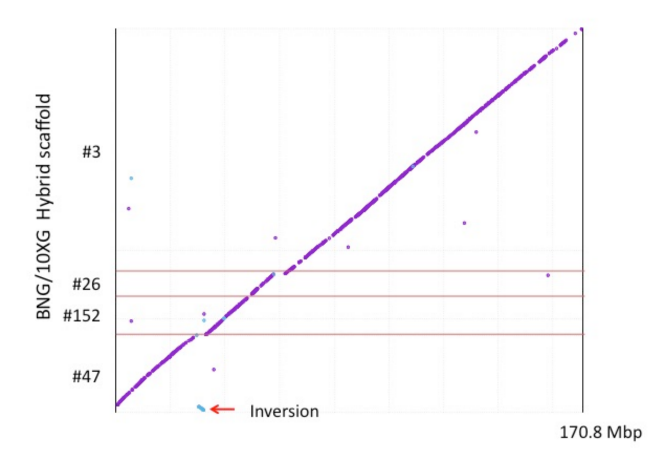


Fig. 5. DNA alignment of the four scaffolds identified by TBLASTN conserved synteny analysis between the monk seal and human chromosome 6 (170.8 Mbp long). Inverted genes in Table 2 are indicated by the arrow.

DISCUSSION:

Next generation sequencing has revolutionized the field of comparative genomics, especially at the level of individual gene sequences. But, creating long assemblies has been less successful because of the complexity of genomes which are frequently interrupted by repeats longer than typical NGS reads. This limitation has led to various innovations such as long-single molecule sequencing (e.g., PacBio, Oxford Nanopore) but throughput, accuracy and costs have limited their widespread adoption. We showed that the 10XG Chromium method, assembled with Supernova v1.1 assembly software, produced long scaffolds 160-fold longer than paired end sequencing assembled with DISCOVAR de novo. When combined with Bionano optical maps the scaffold N50 increased to greater than 29 Mbp, a 200-fold improvement. The technologies described here are a large step toward improving and simplifying de novo genome assembly in that they use orthogonal methods that complement each other. The strengths of the 10XG Chromium Linked-Reads method are that it uses $\sim 10^6$ unique molecular indices to randomly tag long molecules captured in emulsified droplets and then sequences these by standard short-read technology. Because the total amount of DNA is so small (1.2 ng for a mammalian genome), the probability that more than one molecule from the same region of the genome is encapsulated and identically labeled is extremely small, and long molecules will produce linked reads that are likely to span repeats such as L1 and other retrotransposons.

Dong et al. (2013) used optical mapping to assist with the assembly of the goat genome and found that it improved super-scaffold lengths by 5X over those obtained with fosmid end sequencing. Mostovoy et al. (2016) used 10XG chemistry and Bionano Genomics maps to create a phased genome assembly of the widely sequenced human sample NA12878 and reported 170 hybrid scaffolds with an N50 of 33.5 Mbp. We have extended their approach by using newer versions of the 10XG chemistry with an increased number of UMIs and Supernova software for de novo assembly. Although optical maps do not produce sequence per se they do provide a valuable "truth set" of chromosomal structure to confirm or refute sequence assemblies. The Bionano optical maps were useful both for merging 10X scaffolds but also as a quality assessment measure of the overall sequence contiguity in any given scaffold. Although the N50 statistic can be misleading based on the choice of the lower contig cutoff size, the assembly that we produced by combining the 10XG and Bionano optical maps resulted in larger scaffolds than most mammalian genomes reported to date. The hybrid Bionano/10XG scaffolds also had conserved syntenic blocks that were consistent with the human genome and other mammals. The QUAST statistics (Table 1) indicated that 98.3% of the total estimated genome length (i.e., the predicted haploid genome equivalent; 2360/2400 Mbp) was accounted for by the 170 Bionano hybrid scaffolds. This is in agreement with the conserved synteny data which showed that most human chromosome orthologous regions were very well covered. As noted above, BUSCO identified 99% of the conserved gene set within the 170 scaffolds, again indicating that the hybrid scaffolds were of high quality.

The materials cost for this study was less than \$15,000. We anticipate that these costs will become lower in the near term as optical mapping improves, short-read sequencing migrates to higher throughput instruments, or new long read technologies (e.g., nanopore sequencing) mature. Software improvements and more standardized pipelines for genome assembly are also expected to make genome assembly less burdensome.

CONCLUSION:

As a field, comparative genomics is entering a new phase where we expect that the vast majority of living (or recently extinct) species will be sequenced. We know from human studies that many of the regulatory signals that control complex traits and development are in intergenic regions (e.g., Bhatia and Kleinjan, 2014; GWAS Catalog: http://www.ebi.ac.uk/gwas/). Likewise, both the sequences and the genomic architecture of regulatory elements within chromosomes are likely to explain the majority of the phenotypic differences between species. If that is true then a fuller understanding of comparative genomics will require knowing not only selected gene sequences but the context of the genome in which those genes are found.

Combining the two orthogonal methods of Chromium Linked-Reads with Bionano optical maps is likely to make the assembly of high quality genomes routine and significantly improve our understanding of comparative genome biology while reducing costs. Because the Chromium method labels single molecules only a few ng of DNA are needed. This advantage may be particularly useful for non-lethal sampling of organisms both in captivity and in the field. As more species become endangered the need to preserve their genomes for study and conservation will increase. The methods shown here represent a cost-effective and robust strategy to meet that goal.

REFERENCES:

- 1. Anantharaman TS, Mysore V, and B Mishra. 2005. Fast and cheap genome wide haplotype construction via optical mapping. Pac Symp Biocomput 10:385-396.
- 2. Bhatia S and DA Kleinjan 2014. Disruption of long-range gene regulation in human genetic disease. Hum Genet 133:815-45.
- 3. Dong Y, Xie M, Jiang Y, Xiao N, Du X, Zhang W, Tosser-Klopp G, Wang J, Yang S, Liang J, Chen W, Chen J, Zeng P, Hou Y, Bian C, Pan S, Li Y, Liu X, Wang W, Servin B, Sayre B, Zhu B, Sweeney D, Moore R, Nie W, Shen Y, Zhao R, Zhang G, Li J, Faraut T, Womack J, Zhang Y, Kijas J, Cockett N, Xu X, Zhao S, Wang, J and W Wang. 2013. Sequencing and automated whole-genome optical mapping of the genome of a domestic goat (Capra hircus). Nat Biotechnol 31:135-41.
- 4. Gurevich A, Saveliev V, Vyahhi N, and G Tesler. 2013. QUAST: quality assessment tool for genome assemblies. Bioinformatics 29:1072-1075.
- 5. GWAS Catalog. http://www.ebi.ac.uk/gwas/
- 6. Jun J, Mandoiu II, and CE Nelson. 2009. Identification of mammalian orthologs using local synteny. BMC Genomics 10:630.
- 7. Kurtz S, Phillippy A, Delcher AL, Smoot M, Shumway M, Antonescu C and SL Salzberg. 2004. Versatile and open software for comparing large genomes. Genome Biology 5:R12

- 8. Larkin DM, Pape G, Donthu R, Auvil L, Welge M and HA Lewin. 2009. Breakpoint regions and homologous synteny blocks in chromosome have different evolutionary histories. Genome Research 19:770-777.
- 9. Mostovoy Y, Levy-Sakin M, Lam J, Lam ET, Hastie AR, Marks P, Lee J, Chu C, Lin C, Dzakula Z, Cao H, Schlebusch SA, Giorda K, Schnall-Levin M, Wall JD, and PY Kwok. 2016. A hybrid approach for de novo human genome sequence assembly and phasing. Nat. Methods 13:587-590.
- 10. Schultz JK, Baker JD, Toonen RJ, Bowen BW. 2009. Extremely low genetic diversity in the endangered Hawaiian monk seal (Monachus schauinslandi). J Hered. 100:25-33.
- 11. Shelton JM, Coleman MC, Herndon N, Lu N, Lam ET, Anantharaman T, Sheth P, and SJ Brown. 2015. Tools and pipelines for BioNano data: molecule assembly pipeline and FASTA super scaffolding tool. BMC Genomics 16:734.
- 12. Simão FA, Waterhouse RM, Ioannidis P, Kriventseva EV, Zdobnov EM. 2015. BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs. Bioinformatics 31:3210-3212.
- 13. Thorvaldsdottir H, Robinson JT and JP Mesirov. 2012. Integrative Genomics Viewer (IGV): high-performance genomics data visualization and exploration. Brief Bioinform 14:178-92.
- 14. Valouev A, Schwartz DC, Zhou S and MS Waterman. 2006. An algorithm for assembly of ordered restriction maps from single DNA molecules. PNAS 103:15770-75.
- 15. Weisenfeld NI, Yin S, Sharpe T, Lau B, Hegarty R, Holmes L, Sogoloff B, Tabbaa D, Williams L, Russ C, Nusbaum C, Lander ES, MacCallum I, and DB Jaffe. 2014. Comprehensive variation discovery in single human genomes. Nat. Genet. 46:1350-1355.
- 16. Weisenfeld, NI, Kumar, V, Shah, P, Church DM and DB Jaffe. 2016. Direct determination of diploid genome sequences. Bioarchiv preprint http://biorxiv.org/lookup/doi/10.1101/070425
- 17. Xiao M, Phong A, Ha C, Chan TF, Cai D, Leung L, Wan E, Kistler AL, DeRisi JL, Selvin PR, and PY Kwok. 2007. Rapid DNA mapping by fluorescent single molecule detection. Nucleic Acids Res 30:e16.
- 18. Xiao S, Li J, Ma F, Fang L, Xu S, Chen W, and ZY Wang. 2015. Rapid construction of genome map for large yellow croaker (Larimichthys crocea) by the wholegenome mapping in BioNano Genomics Irys system. BMC Genomics 16:670.

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Supplementary materials:

Analysis Commands:

BLAST:

tblastn -num_descriptions 10000 -num_alignments 10000 -evalue 1e-05 -outfmt "7 gacc sacc evalue gstart gend sstart send length score stitle"

BIONANO Hybrid Assembly:

perl /home/bionano/scripts/HybridScaffold/hybridScaffold.pl -n
_fastas__assembly_pseudohap2.1.fasta -b EXP_REFINEFINAL1.cmap -c
/home/bionano/scripts/HybridScaffold/hybridScaffold_config_aggressive.xml -o
./hybridOutput1_noCut -r /home/bionano/tools/RefAligner -B 2 -N 1 -f

BUSCOv2:

BUSCO.py -i reference.fasta -o output_name -l vertebrata -m genome -c 4 -sp human

DISCOVARdenovo:

DiscovarDeNovo READS="*.fastq.gz"NUM_THREADS=48 MAX_MEM_GB=900 OUT_DIR=my_discovar_assembly

NUCMER:

nucmer --maxmatch -c 20 -l 50 reference.fasta query.fasta delta-filter -l 1000 input.delta > out.delta mummerplot -postscript -R reference.fasta -Q query.fasta -f -p out.delta

QUAST:

SUPERNOVA:

supernova run -fastqs=demultiplexed_fastq_path --lanes=1,2
--indices=ATTCCGATA,ATTCCGATC,CCCTAACAA,CCTAACAAT,GAAGGCTGA,TGGATTGCA
--maxreads=975000000 --id=benny --description=monk_seal