Revision: b'bigraph@0.7.0-93-g522fddc'
This file contains statistics about the following genome(s):
$\bullet HG002_HiFi_15kb_37x_hodeco_simple \ HG002_HiFi_15kb_37x \ "retain_cm":"yes","uniquify_ids":"yes","genome":"HG002_HiFi_15kb_37x","quast_mode":"hicanu","read_downsampling_factor":"none","homopolymer_compression":"none","assembler":"hicanu","read_downsampling_factor":"none","homopolymer_compression":"none","assembler":"hicanu","read_downsampling_factor":"none","homopolymer_compression":"none","assembler":"hicanu","read_downsampling_factor":"none","homopolymer_compression":"none","assembler":"hicanu","read_downsampling_factor":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression":"none","homopolymer_compression:"none","homopolymer_com$

Attention: this file was produced automatically, and some statistics might not make sense for certain pipelines.

Table 1: Genome Graph Statistics

Table 2: Algorithm Statistics
Parameter

Table 3: ContigValidator

Table 4: QUAST: # of contigs

Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu		
# contigs (≥ 0 bp)	2,145	2,297	2,913	3,413	3,156	2,059	27,213	33,167	54,391	51,212	54,433		
# contigs (≥ 1000 bp)	2,145	2,297	2,913	3,413	3,156	2,059	27,202	33,167	54,380	51,212	54,433		
# contigs (≥ 5000 bp)	2,111	2,279	2,889	3,334	3,096	2,030	27,103	33,167	30,992	49,828	54,433		
# contigs (> 10000 bp)	1,700	1,964	2,482	2,868	2,719	1,762	22,482	33,161	9,315	48,103	54,429		
# contigs (> 25000 bp)	1,086	1,187	1,537	1,940	1,864	1,103	11,529	23,924	2,769	38,563	27,076		
# contigs (≥ 50000 bp)	825	869	1,116	1,519	1,453	850	5,368	10,984	1,814	25,152	11,827		

Table 5: QUAST: total length of contigs

_ ***** ** ***************************													
Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu		
Total length (≥ 0 bp)	2,710,148,694	2,743,626,278	2,796,824,550	5,360,317,965	5,361,907,129	4,934,073,437	3,586,681,568	5,982,135,458	3,229,928,771	5,714,819,445	6,164,499,418		
Total length (≥ 1000 bp)	2,710,148,694	2,743,626,278	2,796,824,550	5,360,317,965	5,361,907,129	4,934,073,437	3,586,673,578	5,982,135,458	3,229,921,951	5,714,819,445	6,164,499,418		
Total length (≥ 5000 bp)	2,710,004,398	2,743,552,597	2,796,731,921	5,360,023,190	5,361,663,515	4,933,953,564	3,586,333,893	5,982,135,458	3,141,848,600	5,710,415,920	6,164,499,418		
Total length (≥ 10000 bp)	2,707,031,655	2,741,035,875	2,793,458,105	5,356,285,471	5,358,651,204	4,931,807,804	3,549,328,733	5,982,082,078	2,994,888,908	5,697,544,471	6,164,461,200		
Total length (≥ 25000 bp)	2,697,493,053	2,728,625,472	2,778,382,317	5,341,395,358	5,344,725,500	4,921,318,434	3,371,274,581	5,810,363,497	2,902,098,390	5,525,939,457	5,710,291,578		
Total length (≥ 50000 bp)	2,688,295,868	2,717,448,603	2,763,432,540	5,326,413,204	5,330,099,179	4,912,508,287	3,156,598,088	5,386,943,130	2,869,864,868	5,031,848,459	5,215,123,161		

Table 6: QUAST: statistics for contigs > 500bp (or 3000bp for QUAST-LG)

	1 add C O. QUAS1: statistics for contigs ≥ 5000p (or 30000p for QUAS1-LG)													
Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu			
# contigs	2,141	2,296	2,910	3,392	3,150	2,055	27,168	33,167	50,609	50,569	54,433			
Largest contig	71,027,909	74,405,341	44,024,760	62,286,514	62,286,512	149,584,362	36,480,942	143,473,701	31,618,859	45,644,871	132,523,900			
Total length	2,710,138,825	2,743,623,429	2,796,817,553	5,360,269,916	5,361,893,767	4,934,065,233	3,586,601,896	5,982,135,458	3,220,392,699	5,713,400,799	6,164,499,418			
Reference length	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832	3,088,269,832			
GC (%)	40.92	40.86	40.84	40.51	40.51	40.40	40.79	40.86	40.64	40.83	40.85			
Reference GC (%)	40.87	40.87	40.87	40.87	40.87	40.87	40.87	40.87	40.87	40.87	40.87			
N50	14,522,088	13,049,512	9,129,109	16,063,095	16,550,044	32,947,791	2,019,624	2,510,882	3,947,737	260,865	986,873			
NG50	11,774,680	11,087,647	7,770,579	25,164,941	26,493,939	48,303,382	2,623,367	68,318,907	4,158,865	504,342	38,046,558			
N75	5,752,227	5,201,732	3,999,840	7,275,466	8,241,862	13,248,984	412,757	248,560	1,143,928	97,370	152,908			
NG75	3,189,820	3,089,156	2,563,463	18,253,839	18,968,970	34,217,238	1,028,461	32,985,826	1,456,483	342,918	18,172,563			
L50	51	59	89	101	94	47	405	93	211	5,410	309			
LG50	65	73	106	43	40	23	297	16	195	1,727	25			
L75	125	141	205	216	204	112	1,313	3,097	569	14,193	5,132			
LG75	191	204	273	80	74	43	763	32	493	3,601	52			
E-size	19,818,283	17,837,603	11,387,312	19,259,191	20,490,856	37,117,100	3,747,979	35,532,978	5,630,435	1,033,660	23,020,671			

Table 7: QUAST: alignment statistics for contigs ≥ 500bp (or 3000bp for QUAST-LG)

				AST: alignment st							TT. 63
Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu
# misassemblies	1,037	1,571	1,353	3,491	3,723	2,437	10,615	24,550	10,939	21,969	21,346
# unique misassemblies	957	1,450	1,265	2,596	2,782	1,920	7,724	15,559	7,517	12,650	13,400
# misassembled contigs	449	561	551	1,362	1,362	724	1,814	1,307	6,400	1,500	1,560
Misassembled contigs length	1,401,181,134	1,485,206,940	875,538,566	3,809,283,394	4,124,446,179	4,222,179,816	842,057,669	3,102,880,165	1,082,365,816	535,378,803	2,792,253,308
# local misassemblies	2,705	3,077	2,980	6,020	6,173	4,463	14,862	22,082	31,026	21,108	22,421
# unique local misassemblies	2,407	2,788	2,740	3,613	3,691	3,087	6,716	9,630	18,968	9,073	10,042
# scaffold gap ext. mis.	0	0	0	0	0	0	0	0	0	0	0
# scaffold gap loc. mis.	0	0	0	0	0	0	0	0	0	0	0
# possible TEs	166	204	208	436	430	348	1,116	3,768	1,528	2,832	3,226
# unaligned mis. contigs	113	144	151	218	196	154	420	481	1,540	432	525
# unaligned contigs	361 + 719p	570 + 917p	636 + 1037p	780 + 1778p	729 + 1691p	566 + 979p	912 + 3314p	558 + 2297p	11149 + 10522p	442 + 2993p	1040 + 2803p
Unaligned length	19,734,000	25,668,873	24,481,216	43,026,389	43,798,578	30,634,458	139,622,427	191,202,834	134,740,112	176,983,395	200,297,559
Genome fraction (%)	91.39	92.31	94.17	94.97	94.99	94.58	97.78	98.40	95.78	98.18	98.42
Duplication ratio	1.00	1.00	1.00	1.91	1.91	1.76	1.20	2.00	1.10	1.92	2.06
# N's per 100 kbp	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
# mismatches per 100 kbp	118.00	120.91	120.13	131.66	132.10	120.48	202.45	180.88	242.47	178.70	180.52
# indels per 100 kbp	29.67	30.01	29.99	31.29	31.32	30.06	29.68	27.49	187.78	27.59	36.23
Largest alignment	47,921,947	74,389,239	44,016,462	56,946,514	56,946,513	74,384,115	32,046,407	90,496,567	31,574,696	32,050,878	90,221,157
Total aligned length	2,689,347,269	2,716,638,451	2,771,232,004	5,310,842,047	5,311,649,189	4,898,948,974	3,438,541,411	5,771,642,014	3,081,217,872	5,517,535,823	5,945,192,324
EAmeanmax	12,181,695	11,571,860	8,947,360	12,882,900	13,152,452	16,081,291	3,894,055	29,821,065	4,807,323	1,051,676	24,970,912
Strict EAmeanmax	3,405,595	3,122,146	3,031,055	3,401,611	3,425,141	3,433,567	2,256,757	5,564,594	649,471	473,214	5,296,691
EA50max	8,261,576	8,164,963	6,744,390	10,010,056	10,030,572	10,091,829	2,389,466	25,620,191	3,317,710	228,815	21,929,637
Strict EA50max	2,529,467	2,358,674	2,263,458	2,578,478	2,670,188	2,560,581	1,582,183	4,301,869	493,607	219,118	4,167,533
EA75max	2,368,326	2,272,024	2,159,649	2,985,083	3,084,569	3,003,362	895,996	10,224,387	1,077,309	73,548	7,094,515
Strict EA75max	810,871	827,969	841,911	972,310	977,370	963,900	602,162	1,650,652	208,213	71,194	1,526,825
P5k	0.87	0.88	0.90	0.90	0.90	0.90	0.93	0.93	0.91	0.93	0.93
P10k	0.87	0.88	0.89	0.90	0.90	0.90	0.93	0.93	0.90	0.93	0.93
P15k	0.87	0.88	0.89	0.90	0.90	0.90	0.93	0.93	0.90	0.92	0.93
P20k	0.87	0.88	0.89	0.90	0.90	0.90	0.92	0.93	0.90	0.91	0.93
Strict P5k	0.87	0.88	0.90	0.90	0.90	0.90	0.93	0.93	0.91	0.93	0.93
Strict P10k	0.87	0.88	0.89	0.90	0.90	0.90	0.93	0.93	0.90	0.93	0.93
Strict P15k	0.87	0.88	0.89	0.90	0.90	0.90	0.92	0.93	0.90	0.92	0.93
Strict P20k	0.86	0.87	0.89	0.90	0.90	0.90	0.92	0.93	0.89	0.91	0.93
NGA50	8,261,576	8,164,963	6,744,390	17,256,574	17,266,299	29,837,383	2,389,466	25,620,191	3,376,207	428,635	21,929,637
NGA75	2,368,326	2,280,694	2,174,040	13,205,281	13,567,632	16,197,583	897,039	10,224,387	1,165,405	290,138	7,094,515
LGA50	89	101	124	60	58	37	331	36	239	2,165	44
LGA75	252	271	314	111	108	73	855	85	614	4,366	103
EA-size	13,998,976	13,164,487	9,978,229	14,243,069	14,509,266	21,454,715	3,506,248	16,122,283	4,888,781	729,740	13,091,772
	20,000,010	25,101,101	2,910,220	,-10,000	= =,500,200	==,101,110	2,300,210	-5,122,200	2,000,101	. 20,1 10	,501,112

Table 8: QUAST: misassembly statistics for contigs > 500bp (or 3000bp for QUAST-LG)

Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu
# misassemblies	1,037	1,571	1,353	3,491	3,723	2,437	10,615	24,550	10,939	21,969	21,346
# contig misassemblies	1,037	1,571	1,353	3,491	3,723	2,437	10,615	24,550	10,939	21,969	21,346
# c. relocations	799	1,305	1,122	2,562	2,765	1,789	8,352	18,548	9,854	18,021	17,414
# c. translocations	207	230	211	883	910	598	2,205	5,906	1,030	3,868	3,842
# c. inversions	31	36	20	46	48	50	58	96	55	80	90
# scaffold misassemblies	0	0	0	0	0	0	0	0	0	0	0
# s. relocations	0	0	0	0	0	0	0	0	0	0	0
# s. translocations	0	0	0	0	0	0	0	0	0	0	0
# s. inversions	0	0	0	0	0	0	0	0	0	0	0
# misassembled contigs	449	561	551	1,362	1,362	724	1,814	1,307	6,400	1,500	1,560
Misassembled contigs length	1,401,181,134	1,485,206,940	875,538,566	3,809,283,394	4,124,446,179	4,222,179,816	842,057,669	3,102,880,165	1,082,365,816	535,378,803	2,792,253,308
# local misassemblies	2,705	3,077	2,980	6,020	6,173	4,463	14,862	22,082	31,026	21,108	22,421
# scaffold gap ext. mis.	0	0	0	0	0	0	0	0	0	0	0
# scaffold gap loc. mis.	0	0	0	0	0	0	0	0	0	0	0
# possible TEs	166	204	208	436	430	348	1,116	3,768	1,528	2,832	3,226
# unaligned mis. contigs	113	144	151	218	196	154	420	481	1,540	432	525
# mismatches	3,173,515	3,284,752	3,329,042	6,992,265	7,016,906	5,902,152	6,961,350	10,439,981	7,470,963	9,859,716	10,732,429
# indels	798,002	815,128	831,189	1,661,904	1,663,691	1,472,592	1,020,607	1,586,770	5,785,884	1,522,171	$2,\!153,\!717$
# indels ($\leq 5 \text{ bp}$)	670,007	686,089	699,630	1,404,711	1,406,294	1,243,826	831,766	1,276,870	5,066,202	1,230,237	1,838,425
# indels (> 5 bp)	127,995	129,039	131,559	257,193	257,397	228,766	188,841	309,900	719,682	291,934	315,292
Indels length	8,546,640	8,546,771	8,690,018	16,527,152	16,555,050	14,732,079	11,491,876	18,778,818	19,743,048	16,927,204	19,251,051

Table 9: Resource usage

Parameter	w2 noho	w2	w2 sfa	w2 YV sfa	w2 YV	w2 frg YV	flye	hifiasm	mdbg	lja	HiCanu	
time [s]	35,127.96	69,508.90	66,659.66	77,942.09	81,198.64	85,877.10	98,493.00	97,377.00	1,072.00	34,878.00	160,198.00	
mem [GiB]	144.05	136.04	135.93	135.93	136.04	136.04	199.61	155.84	18.40	168.11	77.63	
hoco time		448.05	448.05	448.05	448.05	448.05	N/A	N/A	N/A	N/A	N/A	
hoco mem		0.38	0.38	0.38	0.38	0.38	N/A	N/A	N/A	N/A	N/A	
hodeco time		36,631.00	34,278.00	39,482.00	41,693.00	47,956.00	N/A	N/A	N/A	N/A	N/A	
hodeco mem		1.79	1.79	1.95	1.94	1.94	N/A	N/A	N/A	N/A	N/A	
trivial_omnitigs time				423.67	397.92	0.47	N/A	N/A	N/A	N/A	N/A	
trivial_omnitigs mem				26.77	26.77	0.00	N/A	N/A	N/A	N/A	N/A	
assembly time	29,305.00	27,270.00	26,625.00	26,625.00	27,270.00	27,270.00	98,493.00	97,377.00	1,072.00	34,878.00	160,198.00	
assembly mem	144.05	136.04	135.93	135.93	136.04	136.04	199.61	155.84	18.40	168.11	77.63	
wtdbg2_extract time	851.96	571.85	584.61	584.61	571.85	571.85	N/A	N/A	N/A	N/A	N/A	
wtdbg2_extract mem	0.00	0.00	0.00	0.00	0.00	0.00	N/A	N/A	N/A	N/A	N/A	
contig_assembly time				1,326.76	1,371.82	1,308.73	N/A	N/A	N/A	N/A	N/A	
contig_assembly mem				125.01	125.02	125.02	N/A	N/A	N/A	N/A	N/A	
wtdbg2_consensus time	4,971.00	4,588.00	4,724.00	9,052.00	9,446.00	8,322.00	N/A	N/A	N/A	N/A	N/A	
wtdbg2_consensus mem	6.39	7.41	7.56	9.15	8.86	7.38	N/A	N/A	N/A	N/A	N/A	













































