## Supplementary Figures and Tables

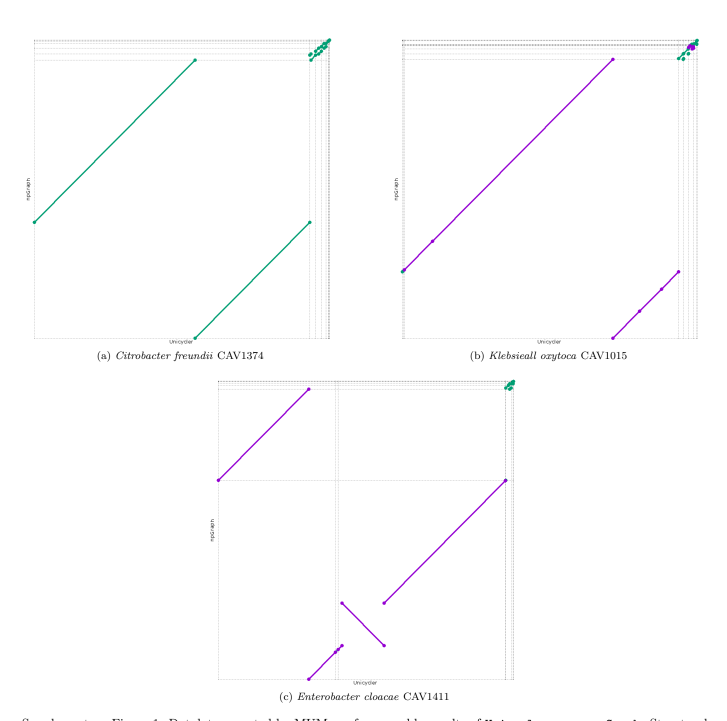
Supplementary Table 1: Benchmarking npGraph against npScarf, hybridSPAdes and Unicycler hybrid assembler with the synthetic data set.

	Aggambler		N50	Mis-	Mismatch	Indel
Method	Assembly size (bp)	#Contigs	(bp)	assemblies	(per 100Kbp)	(per 100Kbp)
random sequences	( - /	#Contigs	(pp)	assembnes	(per 100Kbp)	(per 100Kbp)
npScarf	4110000	3	4000000	0	0.00	0.00
_						
npGraph-bwa	4110000	3	4000000	0	0.00	0.00
npGraph-mm2	4110000	3	4000000	0	0.00	0.00
hybridSPAdes	4110231	3	4000077	0	0.00	0.07
Unicycler	4110000	3	4000000	0	0.00	0.00
random sequences			1000=15		0.00	0.00
npScarf	4110940	3	4002715	0	0.00	0.66
npGraph-bwa	4110000	3	4000000	0	0.00	0.00
npGraph-mm2	4110000	3	4000000	0	0.00	0.00
hybridSPAdes	4107566	3	3999364	0	0.02	0.02
Unicycler	4110000	3	4000000	0	0.00	0.00
random sequences	many repeat					
npScarf	4316934	8	3963485	24	11.55	34.54
npGraph-bwa	4110000	3	4000000	0	0.32	0.15
npGraph-mm2	4110000	3	4000000	0	0.32	0.15
hybridSPAdes	4108190	3	3999621	0	0.68	0.15
Unicycler	4110000	3	4000000	0	0.32	0.15
Acinetobacter A1						
npScarf	3912299	3	3870269	4	4.74	13.02
npGraph-bwa	3918374	2	3909643	0	16.34	0.61
npGraph-mm2	3885898	2	3877167	1	18.05	1.03
hybridSPAdes	3929948	53	3353679	0	35.48	3.22
Unicycler	3917745	2	3909014	0	2.50	0.13
Acinetobacter AB						
npScarf	4512464	7	4304628	35	57.95	72.87
npGraph-bwa	4335342	2	4148952	1	16.93	1.45
npGraph-mm2	4335790	1	4335790	0	6.97	0.25
hybridSPAdes	4337369	3	2701005	0	12.80	1.39
Unicycler	4333041	1	4333041	1	6.42	0.53
E. coli K12 MG16			1000011		0.12	0.00
npScarf	4649902	2	4641702	4	14.94	34.35
npGraph-bwa	4641743	1	4641743	0	4.50	0.43
npGraph-mm2	4641820	1	4641820	0	3.88	0.45 $0.26$
hybridSPAdes	4644555	1	4641036	0	0.62	0.09
Unicycler	4641650	1	4641650	0	3.43	0.26
E. coli O25b H4-S		1	4041000		5.45	0.20
npScarf	5245913	3	5095571	7	7.05	18.81
_	5245913 $5237821$		4049493			0.31
npGraph-bwa		7		1	3.38	
npGraph-mm2	5249799	3	5110117	0	2.40	0.15
hybridSPAdes	5252762	8	4258948	2	5.43	0.57
Unicycler	5249442	3	5109760	0	4.02	0.27
Klebsiella 30660 N		-	FOFOCEO	,		10.10
npScarf	5559772	7	5259053	4	17.18	13.48
npGraph-bwa	5534843	8	5263229	0	3.15	0.76
npGraph-mm2	5534878	8	5263264	0	2.75	0.74
hybridSPAdes	5545668	8	5545668	2	4.95	0.09
Unicycler	5537860	9	5263196	0	1.34	0.51

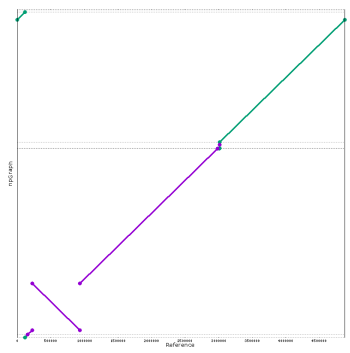
Continued on next page

Supplementary Table 1 - Continued from previous page

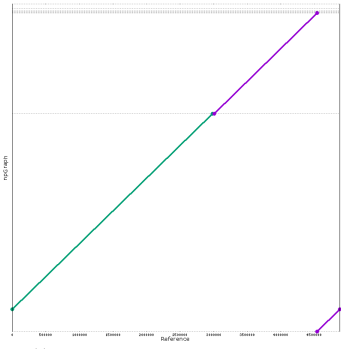
Supplementary Table 1 – Continuea from previous page										
N f - 4 l J	Assembly	// ( + :	N50	Mis-	Mismatch	Indel				
Method	size (bp)	#Contigs	(bp)	assemblies	(per 100Kbp)	(per 100Kbp)				
Klebsiella MGH 78		-	F91.0490	10	14.00	00.07				
npScarf	5729304	5	5316429	12	14.90	20.27				
npGraph-bwa	5695801	7	5311745	1	12.65	1.25				
npGraph-mm2	5696302	6	5315267	0	6.06	0.44				
hybridSPAdes	5706470	11	5315273	1	3.82	0.67				
Unicycler	5694231	14	5315096	0	5.38	0.21				
Klebsiella NTUH-F										
npScarf	5471696	2	5249198	6	4.82	8.55				
npGraph-bwa	5472629	2	5248476	0	2.52	0.31				
npGraph-mm2	5472655	2	5248503	0	1.21	0.26				
hybridSPAdes	5473572	2	5248894	0	0.44	0.15				
Unicycler	5472697	2	5248545	0	2.41	0.35				
Mycobacterium tuberculosis H37Rv										
npScarf	4498245	4	4402238	8	5.15	2.68				
npGraph-bwa	4411406	1	4411406	0	1.88	0.43				
npGraph-mm2	4411532	1	4411532	0	0.68	0.00				
hybridSPAdes	4413942	1	4410519	0	0.75	0.11				
Unicycler	4411538	1	4411538	0	2.22	0.34				
Saccharomyces cere	evisiae S288	<u> </u>								
npScarf	11986800	24	796769	51	62.12	21.46				
npGraph-bwa	11921736	40	913090	3	38.04	1.94				
npGraph-mm2	11920984	38	913198	2	20.66	0.95				
hybridSPAdes	12027533	45	770543	5	32.58	1.94				
Unicycler	11847655	72	909114	0	21.81	1.04				
Shigella dysenteria	e Sd197									
npScarf	-	-	_	_	-	-				
npGraph-bwa	4564058	6	4369264	5	80.64	11.16				
npGraph-mm2	4558920	6	4364264	7	75.51	10.98				
hybridSPAdes	4519131	23	821249	96	9.57	1.42				
Unicycler	4560901	3	4369231	0	11.88	1.05				
Shigella sonnei 53G										
npScarf	6441461	20	1953896	82	164.02	219.52				
npGraph-bwa	5211544	4	4988532	0	14.53	0.31				
npGraph-mm2	5211527	4	4988519	0	8.56	0.17				
hybridSPAdes	5223875	8	2195455	$\overset{\circ}{2}$	41.92	0.06				
Unicycler	5220517	5	4988548	0	7.39	0.52				
Streptococcus suis		0	1000040	0	1.00	0.02				
npScarf	$\frac{2183951}{2183951}$	3	2146594	0	21.51	9.17				
npGraph-bwa	2154623	6	2131479	1	5.25	0.28				
npGraph-bwa npGraph-mm2	2134023		2131479 2146774		5.25 1.44	0.28				
hybridSPAdes	2149870	$\frac{6}{2}$	2146774	$0 \\ 0$	6.82	0.28				
Unicycler  Unicycler	2172703	2	2146257 $2146250$	0	$\frac{0.82}{2.67}$	0.09 $0.32$				
Omeyelei	4170649	Z	2140250	U	2.07	0.52				



Supplementary Figure 1: Dotplot generated by MUMmer for assembly results of Unicycler versus npGraph. Structural agreements between two methods were found in (a) *C.freundii* and (b) *K.oxytoca* assembly contigs. On the other hand, for (c) *E.cloacae* sample, there was a disagreement detected between 2 largest contigs given by two assembly algorithms. This case is investigated more thoroughly by using a reference from a same bacterial strain in Figure 2.



(a)  $\it E.~cloacae$  Unicycler assembly versus reference genome



(b)  $\it E.~cloacae$  npGraph assembly versus reference genome

Supplementary Figure 2: Alignments of an *Enterobacter cloacae* reference genome to assembly sequences generated by (a) Unicycler and (b) npGraph. While the former presents a structural variant, the latter is virtually an 1-to-1 mapping.