

Supplementary Data

Table S1. Summary and accession information for all NCBI reference genomes used in the simulated Illumina sequencing for this study.

See attached file: simulatedDatasets.xlsx

Table S2. Performance of automated binning programs compared to a previous semi-manual approach (Miller, Vanee, et al., 2016) in recovery of the “*Candidatus Endobugula sertula*”.

Method	Length (Mbp)	no. contigs	% "Ca. E. sertula" recovered		completeness	purity
			(contigs, length)			
Autometa	3.73	186	96.7% (101, 3,218,088 bp)		96.2	96.5
MaxBin	3.71	175	91.8% (88, 3,055,541 bp)		96.2	96.0
MetaBAT	2.9	90	86.6% (86, 2,881,097 bp)		93.2	98.0
MyCC	2.84	168	68.7% (73, 2,286,775 bp)		71.6	98.2
semi-manual	3.32	117	100% (117, 3,326,817 bp)		96.2	97.5

Table S3. Percentage of binned sequence by length based on kingdom level classification as inferred from Autometa’s Lowest Common Ancestor (LCA) workflow.

Method	Bacteria	Eukaryota	Viruses	Archaea	unclassified
Autometa	100	0.0	0.0	0.0	0.0
MyCC	24.7	13.5	0.2	0.1	61.5
MaxBin	28.5	11.9	0.2	0.1	59.4
MetaBAT	24.6	13.2	0.2	0.1	61.9

Table S4. Results of MyCC binning on AB1_ovicells contigs that were unfiltered by Autometa's LCA workflow.

MyCC cluster	Size (Mbp)	N ₅₀	No. contigs	CheckM completeness (%)	CheckM purity (%)	Coverage	GC (%)	CheckM lineage
Cluster.8	136.29	9923	16825	41.5	87.5	30.5	33.5	k_Archaea
Cluster.2	31.68	14997	2811	37.6	77.3	2.1	34.5	k_Archaea
Cluster.16	17.12	11618	1879	74.4	65.1	5.0	41.3	k_Archaea
Cluster.5	11.61	9846	1474	89.6	82.9	1.6	41.2	k_Bacteria
Cluster.1	7.76	33113	430	93.8	89.8	2.3	50.4	c_Gammaproteobacteria
Cluster.19	4.17	46272	231	92.6	92.7	2.0	60.4	f_Rhodobacteraceae
Cluster.14	3.45	20489	283	87.9	54.9	5.0	46.4	k_Bacteria
Cluster.11	3.42	6074	561	40.0	88.3	1.6	53.9	f_Rhodobacteraceae
Cluster.12	3.13	10960	354	24.1	94.8	4.2	43.8	k_Bacteria
Cluster.20	3.00	4960	609	34.0	98.8	1.3	50.8	k_Bacteria
Cluster.4	2.83	43116	168	71.6	98.2	4.4	40.2	c_Gammaproteobacteria
Cluster.18	2.65	6118	406	56.5	89.3	1.7	43.6	c_Gammaproteobacteria
Cluster.13	1.99	27349	126	98.7	100.0	26.6	34.3	k_Bacteria
Cluster.15	1.78	6310	303	67.9	97.7	1.5	40.1	c_Gammaproteobacteria
Cluster.22	1.31	4099	299	26.0	100.0	6.2	33.1	s_algicola
Cluster.10	1.06	4363	233	40.3	100.0	1.3	48.9	c_Gammaproteobacteria
Cluster.9	0.83	4168	195	16.7	100.0	1.2	49.3	root
Cluster.7	0.81	4491	172	17.8	99.6	1.4	49.3	c_Gammaproteobacteria
Cluster.6	0.72	4047	172	17.2	100.0	1.2	48.2	k_Bacteria
Cluster.3	0.70	593222	16	15.7	100.0	16.4	22.8	k_Bacteria
Cluster.21	0.65	17313	58	61.5	99.8	6.7	23.7	k_Bacteria
Cluster.17	0.60	16076	65	10.3	94.0	7.8	33.3	p_Cyanobacteria

Table S5. Length of sequence (in bp) based on kingdom level classification in MyCC bins that were unfiltered by Autometa's LCA workflow.

MyCC_cluster	Bacteria	Eukaryota	Viruses	Archaea	unclassified	total length
Cluster.8	3,129,421	23,629,421	38,514	299,748	109,189,222	136,286,326
Cluster.2	1,763,368	4,663,853	29,801	34,257	25,190,411	31,681,690
Cluster.16	2,480,635	3,342,491	61,185	40,085	11,198,545	17,122,941
Cluster.5	11,555,610	0	0	0	52,497	11,608,107
Cluster.1	7,702,631	5,550	0	0	51,469	7,759,650
Cluster.19	4,171,828	0	0	0	0	4,171,828
Cluster.14	3,413,645	0	0	11,925	28,344	3,453,914
Cluster.11	3,373,766	9,037	0	0	34,513	3,417,316
Cluster.12	3,067,560	3,490	0	28,361	33,458	3,132,869
Cluster.20	2,935,365	5,355	0	0	54,441	2,995,161
Cluster.4	2,786,399	8,449	0	3,288	34,683	2,832,819
Cluster.18	2,648,425	0	0	0	3,763	2,652,188
Cluster.13	1,777,789	140,579	0	0	72,184	1,990,552
Cluster.15	1,770,202	0	0	0	8,513	1,778,715
Cluster.22	1,254,005	15,479	0	27,945	11,970	1,309,399
Cluster.10	1,055,973	0	0	0	0	1,055,973
Cluster.9	829,401	0	0	0	0	829,401
Cluster.7	785,921	0	0	11,309	16,013	813,243
Cluster.6	718,005	0	0	0	0	718,005
Cluster.3	608,007	19,179	0	0	73,961	701,147
Cluster.21	538,243	35,539	0	0	78,722	652,504
Cluster.17	290,066	218,709	0	4,042	88,344	601,161

Table S6. Fraction of reference genome in MIX-51 successfully assembled by metaSPAdes (v3.9.0) (Nurk *et al.*, 2017) and F1 for each binning program as determined by alignments to known reference genomes using metaQUAST (Mikheenko *et al.*, 2016).

Genome	Fraction Assembled (%)	Autometa F1	MyCC F1	MaxBin F1	MetaBAT F1
<i>Alistipes indistinctus</i>	99.5	0.996	0.989	0.966	0.996
<i>Bacteroides cellulosilyticus</i>	93.1	0.859	0.276	0.139	0.345
<i>Bacteroides dorei</i>	83.3	0.000	0.181	0.415	0.158
<i>Bacteroides finegoldii</i>	91.9	0.000	0.180	0.098	0.192
<i>Bacteroides intestinalis</i>	94.7	0.904	0.244	0.244	0.288
<i>Bacteroides ovatus</i>	95.6	0.639	0.260	0.317	0.283
<i>Bacteroides plebeius</i>	95.4	0.928	0.930	0.227	0.175
<i>Bacteroides stercoris</i>	95.4	0.915	0.449	0.388	0.505
<i>Bacteroides thetaiotaomicron</i> 3731	59.9	0.000	0.113	0.261	0.204
<i>Bacteroides thetaiotaomicron</i> 7330	68.2	0.000	0.095	0.366	0.264
<i>Bacteroides thetaiotaomicron</i> VPI-5482	78.1	0.022	0.148	0.547	0.390
<i>Bacteroides uniformis</i>	96.2	0.981	0.686	0.588	0.877
<i>Bacteroides vulgatus</i>	66.3	0.000	0.130	0.320	0.110
<i>Bifidobacterium adolescentis</i>	95.4	0.790	0.399	0.443	0.523
<i>Bifidobacterium angulatum</i>	97.3	0.736	0.347	0.672	0.349
<i>Bifidobacterium bifidum</i>	98.9	0.966	0.961	0.961	0.966
<i>Bifidobacterium dentium</i>	97.9	0.566	0.450	0.480	0.578
<i>Bifidobacterium pseudocatenulatum</i>	89.6	0.525	0.319	0.806	0.723
<i>Blautia hansenii</i>	96.5	0.947	0.516	0.904	0.957
<i>Blautia luti</i> DSM 14534	68.0	0.724	0.314	0.376	0.712
<i>Citrobacter youngae</i>	99.1	0.897	0.449	0.575	0.540
<i>Clostridium asparagiforme</i>	96.7	0.974	0.977	0.975	0.972
<i>Clostridium bolteae</i>	97.3	0.894	0.493	0.968	0.963
<i>Clostridium hathewayi</i>	88.3	0.925	0.507	0.454	0.765
<i>Clostridium hylemonae</i>	97.5	0.947	0.947	0.947	0.947
<i>Clostridium M62</i> 1	93.8	0.941	0.931	0.935	0.917
<i>Clostridium nexile</i>	89.5	0.829	0.800	0.408	0.564
<i>Clostridium ramosum</i>	99.2	0.989	0.571	0.807	0.558
<i>Clostridium sporogenes</i>	99.0	1.000	0.990	0.995	1.000
<i>Clostridium symbiosum</i>	98.8	0.929	0.412	0.670	0.367

<i>Collinsella intestinalis</i>	98.9	0.999	0.601	0.999	0.930
<i>Collinsella stercoris</i>	94.0	0.795	0.551	0.792	0.746
<i>Coprococcus comes</i>	95.9	0.844	0.439	0.204	0.624
<i>Dorea formicigenerans</i>	94.8	0.868	0.432	0.210	0.682
<i>Edwardsiella tarda</i>	96.1	0.951	0.951	0.951	0.772
<i>Enterobacter cancerogenus</i>	98.5	0.998	0.473	0.763	0.561
<i>Escherichia fergusonii</i>	97.1	0.961	0.473	0.968	0.694
<i>Eubacterium biforme</i>	94.4	0.985	0.973	0.901	0.903
<i>Eubacterium eligens</i>	98.8	0.988	0.983	0.981	0.986
<i>Holdemania filiformis</i>	95.9	0.989	0.982	0.948	0.982
<i>Lactobacillus reuteri</i>	95.2	0.988	0.978	0.978	0.975
<i>Lactobacillus ruminis</i>	82.6	0.988	0.988	0.988	0.899
<i>Marvinbryantia formatexigens</i>	98.6	0.983	0.965	0.943	0.961
<i>Megamonas funiformis</i>	97.4	0.999	0.994	0.789	0.999
<i>Parabacteroides johnsonii</i>	91.7	0.493	0.605	0.555	0.547
<i>Parabacteroides merdae</i>	93.8	0.570	0.628	0.574	0.662
<i>Proteus penneri</i>	99.5	0.993	0.990	0.988	0.717
<i>Roseburia intestinalis</i>	93.4	0.957	0.904	0.908	0.940
<i>Ruminococcus gnavus</i>	95.2	0.797	0.821	0.172	0.433
<i>Streptococcus infantarius</i>	99.0	0.999	0.985	0.992	0.754
<i>Subdoligranulum variabile</i>	97.0	0.626	0.691	0.957	0.593

Table S7. Recall and precision values for *Bacteroides thetaiotaomicron* strains in Mix-51.

		Autometa	MyCC	MaxBin	MetaBAT
recall		0	94	62	59
precision		0	5	26	17
<i>B. thetaiotaomicron</i> 7330	recall	3	95	70	64
	precision	2	8	45	28
<i>B. thetaiotaomicron</i> VPI-5482	recall	0	93	37	38
	precision	0	6	20	14
<i>B. thetaiotaomicron</i> 3731	recall	0	93	37	38
	precision	0	6	20	14

Table S8. Total number of bins recovered by tested binning programs, where “NA” denotes cases where programs failed to complete successfully.

Method	78.125 Mbp	156.25 Mbp	312.5 Mbp	625 Mbp	1250 Mbp	2500 Mbp	5000 Mbp	10000 Mbp
Autometa	23	46	90	171	384	613	940	161
MyCC	23	42	81	139	275	315	NA	NA
MaxBin	22	38	62	109	185	298	565	116
MetaBAT	21	42	75	118	233	347	NA	NA

Table S9. Quantitative data for median F1 values, where “NA” denotes cases where programs failed to complete successfully.

Method	78.125 Mbp	156.25 Mbp	312.5 Mbp	625 Mbp	1250 Mbp	2500 Mbp	5000 Mbp	10000 Mbp
Autometa	1.00	0.98	0.99	0.98	0.98	0.95	0.90	0.01
MyCC	0.99	0.97	0.93	0.88	0.74	0.46	NA	NA
MaxBin	0.98	0.94	0.91	0.67	0.42	0.17	0.23	0.02
MetaBAT	0.97	0.87	0.96	0.60	0.45	0.24	NA	NA

Table S10. Quantitative data for F1 recovery values, where “NA” denotes cases where programs failed to complete successfully.

Method	78.125 Mbp	156.25 Mbp	312.5 Mbp	625 Mbp	1250 Mbp	2500 Mbp	5000 Mbp	10000 Mbp
Autometa	0.98	0.91	0.91	0.89	0.87	0.77	0.60	0.08
MyCC	0.98	0.89	0.92	0.78	0.70	0.48	NA	NA
MaxBin	0.86	0.79	0.66	0.55	0.46	0.37	0.33	0.04
MetaBAT	1.00	0.72	0.77	0.58	0.49	0.40	NA	NA

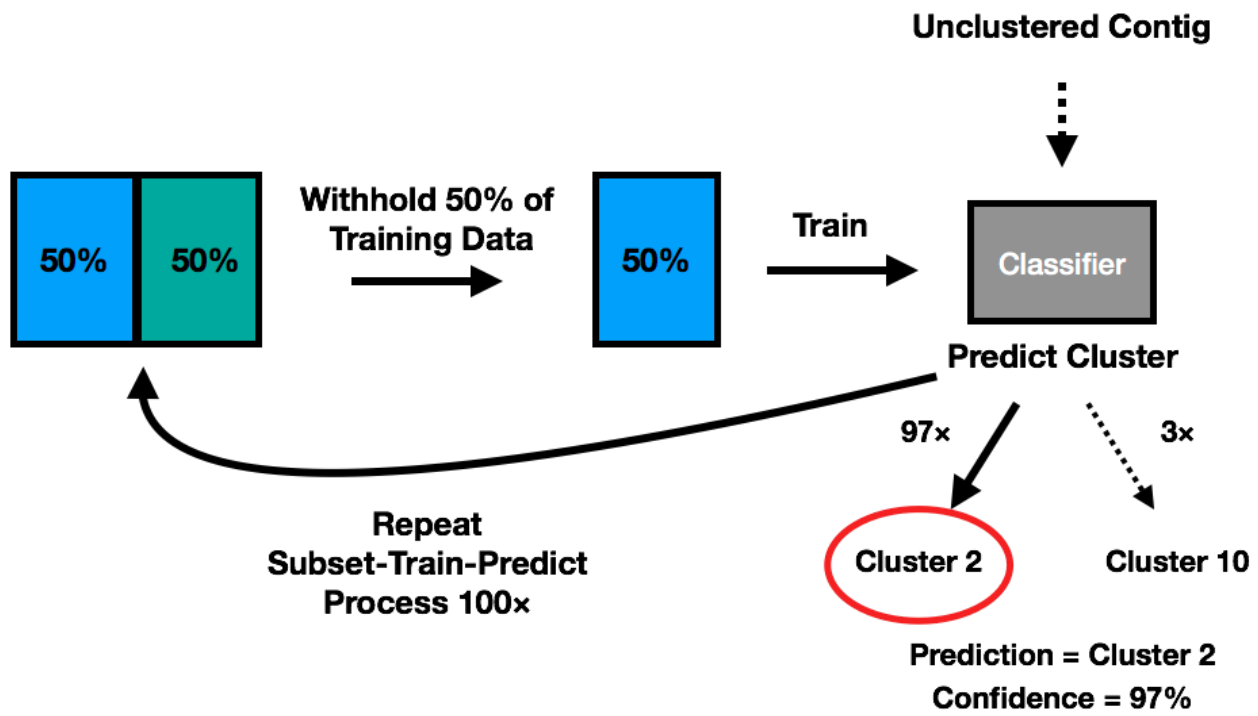


Figure S1. Schematic of jackknife cross validation concept, adapted from (Chevrette *et al.*, 2017).

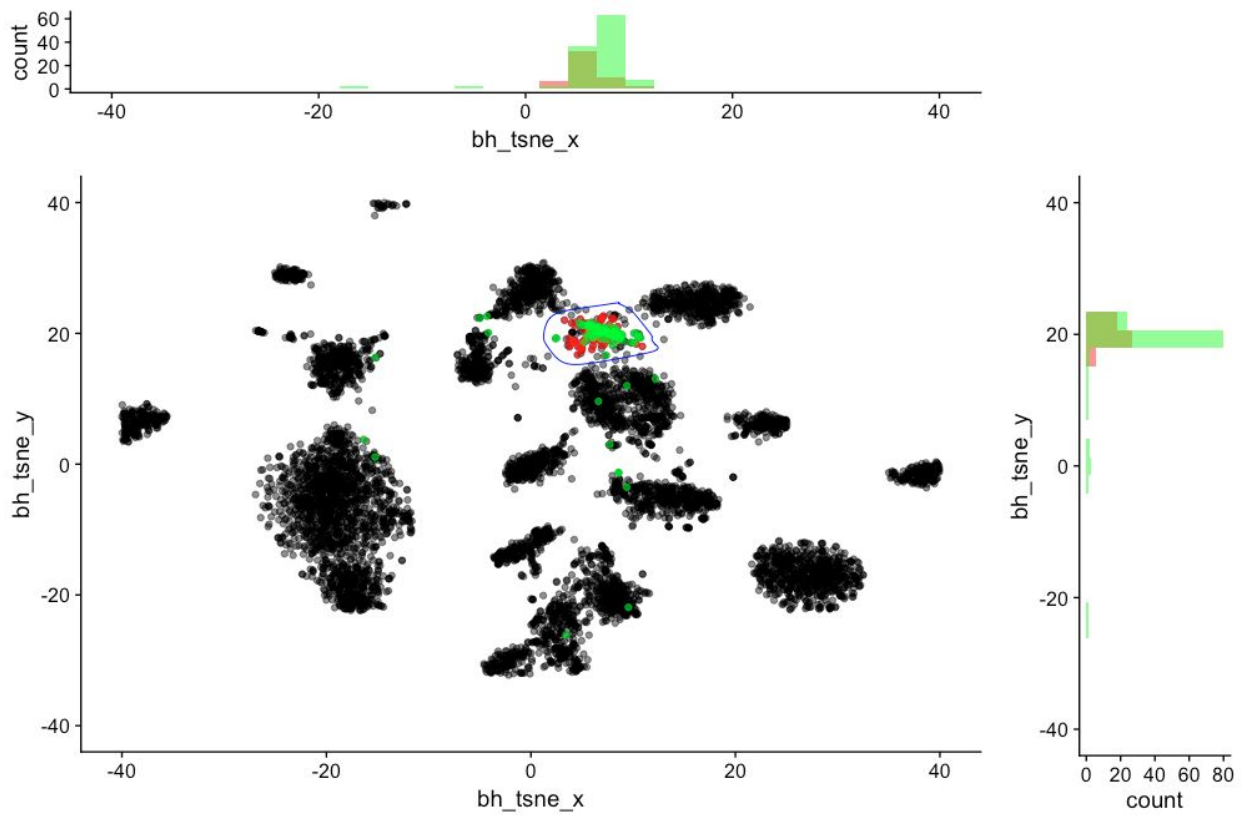


Figure S2. Nucleotide composition profiles of potential “*Ca. E. sertula*” contigs as identified by Autometa and MyCC. Contigs that were newly identified by both Autometa and MyCC as belonging to their “*Ca. E. sertula*” bin are colored in red whereas those that were previously classified as “*Ca. E. sertula*” by our previous semi-manual approach (Miller, Vanee, *et al.*, 2016; Miller, Weyna, *et al.*, 2016) are colored in green.

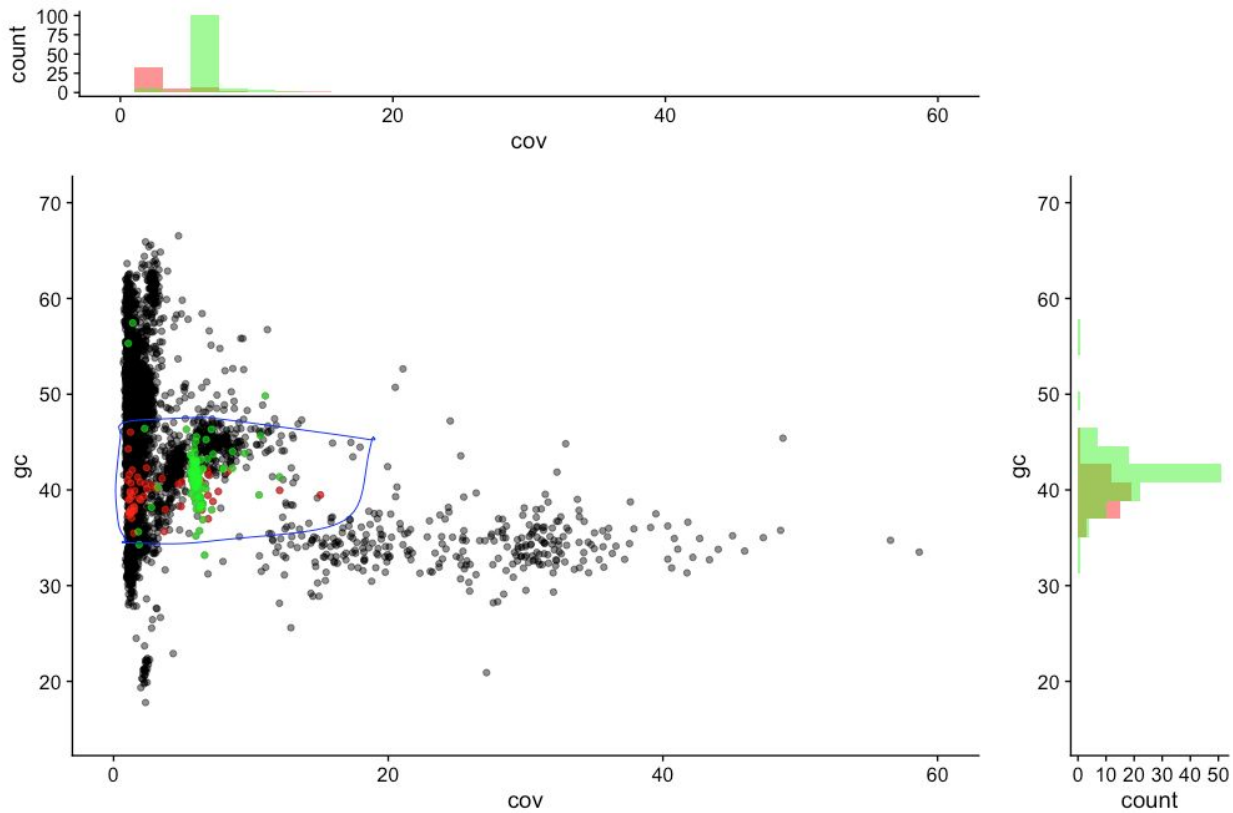


Figure S3. GC% and sequence coverage profiles of potential “*Ca. E. sertula*” contigs as identified by Autometa and MyCC. Contigs that were newly identified by both Autometa and MyCC as belonging to their “*Ca. E. sertula*” bin are colored in red whereas those that were previously classified as “*Ca. E. sertula*” by our previous semi-manual approach (Miller, Vanee, et al., 2016; Miller, Weyna, et al., 2016) are colored in green.

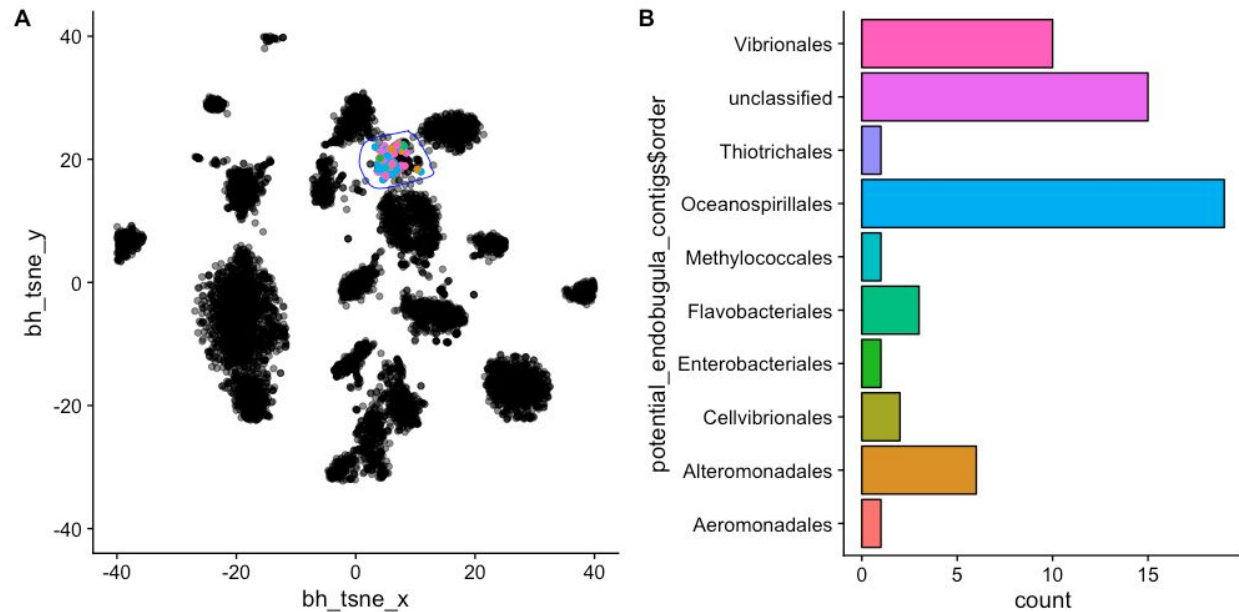


Figure S4. Taxonomic classification of potential “*Ca. E. sertula*” contigs as defined by Autometa’s LCA workflow. Nineteen of 51 contigs are classified as “Oceanospirillales” as the order level. Sixteen of these 19 contigs are classified as “Endozoicomonas” at the genus level and thus likely represent contamination with the “AB1_endozoicomonas genome” bin we identified in a previous study by semi-manual binning approach (Miller, Weyna, et al., 2016).

Supplementary References

- Chevrette,M.G. *et al.* (2017) SANDPUMA: Ensemble Predictions of Nonribosomal Peptide Chemistry Reveals Biosynthetic Diversity across Actinobacteria. *Bioinformatics*.
- Mikheenko,A. *et al.* (2016) MetaQUAST: evaluation of metagenome assemblies. *Bioinformatics*, **32**, 1088–1090.
- Miller,I.J., Vanee,N., *et al.* (2016) Lack of overt genome reduction in the bryostatin-producing bryozoan symbiont ‘Candidatus Endobugula sertula’. *Appl. Environ. Microbiol.*, **82**, 6573–6583.
- Miller,I.J., Weyna,T.R., *et al.* (2016) Single sample resolution of rare microbial dark matter in a marine invertebrate metagenome. *Sci. Rep.*, **6**, 34362.
- Nurk,S. *et al.* (2017) metaSPAdes: a new versatile metagenomic assembler. *Genome Res.*