# CCQM Microbial Identity 16S rRNA Interlaboratory Study

Supplemental Results

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### 1 Biologically Conserved Positions

None of the variants for the biologically conserved positions were called using both variant callers, indicating that the variants were potential false positives (Manuscript Table 2, Tables S4 and S5). Consensus base quality statistics for biologically conserved positions are summarized below (Table S1).

Table S1: Biologically Conserved Position Base Qualities Characteristics of consensus based calls for conserved bases. Normalized quality values were obtained by dividing raw quality score (Raw Qual) assigned by GATK for each biologically conserved base position by the depth of coverage for that position

Org	Plat	Lab	Rep	Raw Qual	Normalized	Min	Max
Ecoli	454	LGC	1	140738.23	2.85	1.25	3.00
Ecoli	454	LGC	2	68081.73	2.85	0.63	2.98
Ecoli	454	LGC	3	128788.23	2.93	1.16	2.99
Ecoli	454	NMIA	1	11457.23	2.51	0.31	2.97
Ecoli	ION	NIMC	1	1165.23	2.78	0.59	3.14
Ecoli	ION	NIST	1	1112.23	2.48	0.51	3.16
Ecoli	Sanger	ATCC	1	34.23	17.11	9.31	31.24
Ecoli	Sanger	ISP	1	31.24	31.23	-10.00	31.24
Ecoli	Sanger	LGC	1	169.23	3.60	0.51	3.97
Ecoli	Sanger	NIST	1	115.23	3.97	-1.43	10.06
Lmono	454	LGC	1	11757.73	1.72	0.52	2.84
Lmono	454	LGC	2	115365.73	2.89	1.43	3.00
Lmono	454	LGC	3	103741.23	2.87	1.44	3.00
Lmono	454	NMIA	1	11635.23	2.41	0.79	2.92
Lmono	ION	NIMC	1	1173.23	2.81	0.33	3.14
Lmono	ION	NIST	1	1265.23	2.56	0.23	2.90
Lmono	Sanger	ATCC	1	34.23	17.11	-10.00	31.24
Lmono	Sanger	ISP	1	34.23	17.11	-10.00	31.24
Lmono	Sanger	LGC	1	169.23	3.45	1.26	3.71
Lmono	Sanger	NIST	1	242.23	3.41	2.18	3.78

All variant calls for the biologically conserved positions were evaluated for being potential false positives Tables S4 and S5. The potential variants identified by the eight variant calling pipelines were analyzed for potential reasons for a false positive variant call. The Fisher Strand bias statistic was used to classify false positive variants due to strand bias (FS; 60). Variants present in non-target regions and at the end of the reference sequence were identified based on positions relative to the reference. False positive variants due to homopolymer systemic sequencing errors and a high proportion of bases covering the identified variant position were identified by visually inspecting the mapping file. Additionally, the contaminant reads were first identified when upon visual inspection of the mapping files revealed a small proportion to highly similar reads that were responsible for a number of variant calls. Note that for the NIST Ion Torrent L. monocytogenes dataset at position 792 a variant was called by the UnifiedGenotyper Variant Calling Algorithm when the reads were mapped using both bwa and tmap, but the FS score was only above 60 when the reads were mapped with tmap. Upon manual inspection of the results we attributed the false positive to a strand bias.

## 2 Biologically Variable Positions

To determine the variant copy ratios, a novel Bayesian analysis based on binomial sampling theory was developed (Supplemental Computational Methods). According to the binomial distribution, the observed variant ratios, while precise (due to high coverage), differed significantly from all potential variant copy ratios. Subsequently given, the observed variant copy ratios, a Bayesian approach was used to identify the most probable variant copy ratio out of the possible ratios assuming *E. coli* and *L. monocytogenes* have seven and six 16S gene copies respectively (Figs. S1 and S2).

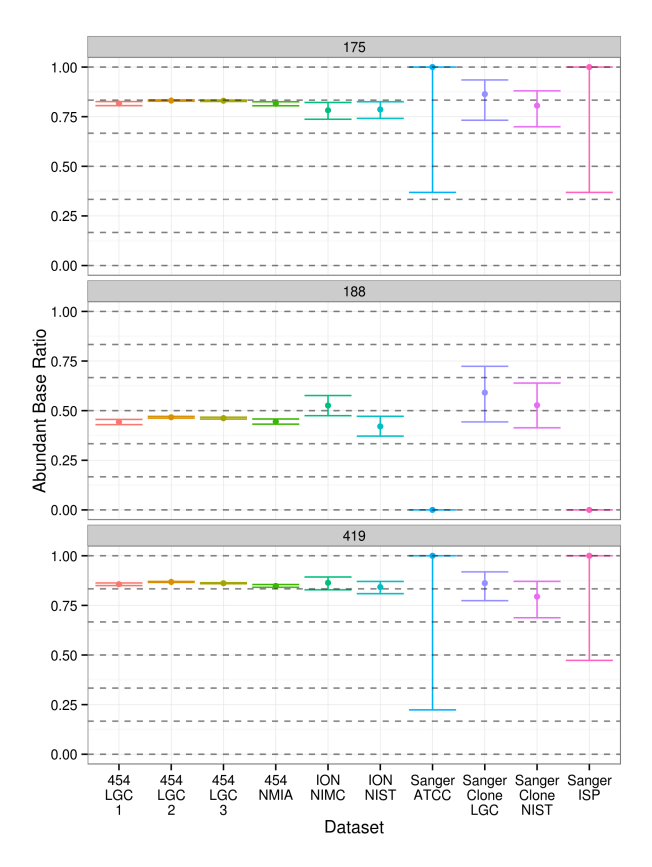


Figure S1: Variant copy ratios at three biologically variable positions (175, 188 and 419) in L. monocytogenes. Variable positions shown in grey box above each graph. Error bars represent the 95 % posterior credibility interval estimated from a beta binomial distribution where is the major variant count + 1 and is the minor variant count + 1. One sided credibile intervals were calcualted for prior probabilities of 0 and 1. Grey dashed lines indicate the potential variant copy ratios assuming six gene copies (i.e. 0:6 corresponds to 0, 2:4 to 0.33, 3:3 to 0.5, 4:2 to 0.66, 5:1 to 0.83 and 6:0 to 1).

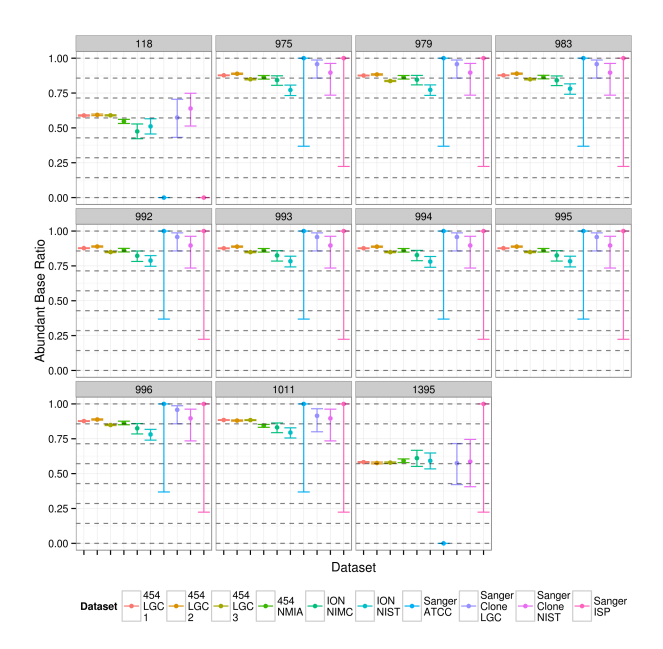


Figure S2: Variant copy ratios at eleven biologically variable positions in  $E.\ coli$ . Variable positions shown in grey box above each graph. Error bars represent the 95 % posterior credibility interval estimated from a beta binomial distribution where is the major variant count + 1 and is the minor variant count + 1. One sided credibile intervals were calcualted for prior probabilities of 0 and 1. Grey dashed lines indicate the potential variant ratios assuming seven gene copies, (i.e. 0:7 to 0; 1:6 to 0.14; 2:5 to 0.26; 3:4 to 0.43, 4:3 to 0.57; 5:2 to 0.71; 6:1 to 0.86; and 7:0 to 1).

## 3 Likely sets of variant combinations

Most likely combination of variant strings for "454" and Sanger Clone library datasets (Table S2 and Table S3).

Table S2: Estimated most likely set of variant combinations for *E. coli*. See supplemental compu-

tation methods for how chimera and likelihood were calculated.

dataset	likelihood	chimera	ACCGATTGTA	ACCGATTGTG	GGTAGAATCA
Ecoli-454-LGC-1	0.04	275.55	3	3	1
Ecoli-454-LGC-2	0.03	275.27	3	3	1
Ecoli-454-LGC-3	0.04	242.13	3	3	1
Ecoli-454-NMIA-1	0.06	30.74	3	3	1
Ecoli-LGC-Sanger-Clones.csv	0.05	3.55	3	4	0
Ecoli-NIST-Sanger-Clones.csv	0.12	4.54	3	4	0
Consensus	0.04	717.62	3	3	1

Table S3: Estimated most likely set of variant combinations for *L. monocyotogenes*. See supple-

mental computation methods for how chimera and likelihood were calculated.

dataset	likelihood	chimera	GCG	GTA	GTG	TCG
Lmono-454-LGC-1	0.00	47.31	2	1	2	1
Lmono-454-LGC-2	0.01	572.64	2	1	2	1
Lmono-454-LGC-3	0.01	319.50	2	1	2	1
Lmono-454-NMIA-1	0.00	55.95	2	1	2	1
Lmono-LGC-Sanger-Clones.csv	0.00	5.13	2	1	2	1
Lmono-NIST-Sanger-Clones.csv	0.01	8.38	2	1	2	1
Consensus	0.01	850.46	2	1	2	1

## 4 Appendix

### Full List of False Positive Variants

All variants called by the 8 piplines used during the pipeline validation along with the suspected cause of the variant. The following abbreviations were used in Tables S4 and S5: Org - Organism, Plat - sequencing platform, Rep - replicate, Map - read mapping algorithm, Var - variant calling algorithm, POS - base position relative to the reference, DP - coverage, QUAL - confidence in variant call assigned my variant calling algorithm, MQ - mapping quality score assigned by mapping algorithm, FS - fisher strain bias test statistic, Cause - hypothesized cause of false positive variant call. See supplemental manuscript methods section for mapping algorithm and variant calling algorithm descriptions.

Table S4: *E. coli* Pipeline Comparison Characteristics of variant calls for different bioinformatic pipelines.

Org	Plat	Lab	Rep	Map	Var	POS	DP	QUAL	MQ	FS	Cause
Ecoli	454	LGC	1	bwa	gatk	324	250	443.77	60.00	47.88	End of read
Ecoli	454	LGC	1	TMAP	gatk	324	250	432.77	88.54	60.26	End of read
Ecoli	454	LGC	1	bwa	gatk	325	250	308.77	60.00	53.48	End of read
Ecoli	454	LGC	1	TMAP	gatk	325	250	309.77	88.54	50.67	End of read
Ecoli	454	LGC	1	bwa	$\operatorname{sam}$	396	2551	81.00	60.00		End of read
Ecoli	454	LGC	1	TMAP	$\operatorname{sam}$	396	3013	37.00	56.00		End of read
Ecoli	454	LGC	1	bwa	gatk	940	19	215.77	60.00	28.54	Non-target region
Ecoli	454	LGC	1	TMAP	gatk	940	21	179.77	80.15	28.54	Non-target region
Ecoli	454	LGC	1	bwa	gatk	959	250	1222.77	60.00	9.12	End of read
Ecoli	454	LGC	2	bwa	gatk	106	250	235.77	60.00	0.00	End of read
Ecoli	454	LGC	2	TMAP	gatk	106	250	34.77	68.90	0.00	End of read
Ecoli	454	LGC	2	bwa	gatk	959	250	795.77	59.98	28.04	End of read
Ecoli	454	LGC	3	bwa	gatk	324	250	231.77	59.83	40.63	End of read
Ecoli	454	LGC	3	TMAP	gatk	324	250	739.77	88.54	63.25	End of read
Ecoli	454	LGC	3	bwa	gatk	325	250	556.77	59.83	60.23	End of read
Ecoli	454	LGC	3	TMAP	gatk	325	250	498.77	88.54	36.85	End of read
Ecoli	454	LGC	3	bwa	gatk	348	250	741.77	59.92	11.62	End of read
Ecoli	454	LGC	3	bwa	$\operatorname{sam}$	417	1032	22.00	60.00		Homopolymer
Ecoli	454	LGC	3	TMAP	$\operatorname{sam}$	417	1020	32.00	58.00		Homopolymer
Ecoli	454	LGC	3	bwa	gatk	940	9	91.05	60.00	0.00	Non-target region
Ecoli	454	LGC	3	TMAP	gatk	940	14	194.29	82.41	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	313	250	5630.77	80.26	453.68	Strand bias
Ecoli	454	NMIA	1	TMAP	gatk	508	250	1160.77	83.71	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	509	250	1208.77	83.71	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	510	250	1275.77	83.71	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	514	250	1185.77	83.71	0.00	End of read
Ecoli	454	NMIA	1	TMAP	$\operatorname{sam}$	514	6337	5.46	59.00		End of read
Ecoli	454	NMIA	1	TMAP	$\operatorname{gatk}$	901	208	8061.77	84.22	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{gatk}$	904	208	8023.77	84.22	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{gatk}$	934	250	8711.77	71.76	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{gatk}$	935	250	8708.77	71.76	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	938	250	8620.77	71.71	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{sam}$	938	2747	9.54	56.00		Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	939	250	8619.77	71.71	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{sam}$	939	2747	15.20	60.00		Non-target region
Ecoli	454	NMIA	1	TMAP	$\operatorname{sam}$	941	2747	9.52	55.00		Non-target region
Ecoli	ION	NIMC	1	TMAP	$\operatorname{sam}$	1463	169	22.50	60.00		End of reference
Ecoli	Sanger	NIST	1	TMAP	$\operatorname{sam}$	1463	29	139.00	60.00		End of reference

Table S5:  $L.\ monocytogenes$  Positions Pipeline Comparison Characteristics of variant calls for different bioinformatic pipelines.

Org	Plat	Lab	Rep	Map	Var	POS	DP	QUAL	MQ	FS	Cause
Lmono	454	LGC	1	bwa	gatk	315	250	4752.77	45.92	101.16	Strand bias
Lmono	454	LGC	1	bwa	gatk	328	250	4865.77	45.96	107.67	Strand bias
Lmono	454	LGC	1	TMAP	$\operatorname{gatk}$	334	250	4700.77	68.98	300.59	Strand bias
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	354	250	62.77	57.95	38.10	End of read
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	366	248	47.77	57.94	40.36	End of read
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	508	250	1386.77	51.13	7.03	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	508	7744	10.40	55.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	533	166	1407.77	46.04	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	533	1763	156.00	38.00		Contaminants
Lmono	454	LGC	1	TMAP	gatk	533	250	1768.77	48.73	0.00	Contaminants
Lmono	454	LGC	1	TMAP	$\operatorname{gatk}$	536	250	1779.77	48.73	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	537	166	1394.77	46.04	1.78	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	537	1763	88.00	38.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	538	166	1634.77	46.04	0.83	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	538	1623	128.00	38.00		Contaminants
Lmono	454	LGC	1	TMAP	$\operatorname{gatk}$	539	250	1798.77	48.73	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	548	166	2211.77	46.04	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	548	1762	201.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	549	166	2217.77	46.04	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	549	1763	186.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	550	166	2247.77	46.04	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	550	1763	175.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	555	167	2077.77	46.14	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	555	1764	222.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	559	167	2201.77	46.14	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	559	1763	222.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	574	168	2288.77	46.03	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	574	1765	189.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	585	168	1737.77	46.03	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	585	1736	213.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	587	168	1980.77	46.03	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	587	1741	212.00	37.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	595	168	2352.77	46.03	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	595	1741	188.00	38.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	677	250	5127.77	58.81	4.08	Contaminants
Lmono	454	LGC	1	bwa	sam	677	4525	222.00	58.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	700	249	5372.77	56.79	14.63	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	700	4604	222.00	58.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	703	249	4820.77	56.79	4.37	Contaminants
Lmono	454	LGC	1	bwa	sam	703	4604	222.00	58.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	712	249	4896.77	56.79	13.22	Contaminants
Lmono	454	LGC	1	bwa	sam	712	4602	222.00	58.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	716	249	4009.77	56.79	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	716	4602	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	729	226	3856.77	58.93	0.72	Contaminants
Lmono	454	LGC	1	bwa	sam	729	4510	222.00	59.00		Contaminants

Lmono	454	LGC	1	bwa	gatk	731	226	2873.77	58.93	0.00	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	$\operatorname{sam}$	731	4487	201.00	59.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	733	226	3028.77	58.93	0.00	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	$\operatorname{sam}$	733	4511	182.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	738	226	4461.77	58.93	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	$\operatorname{sam}$	738	4493	222.00	59.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	$\operatorname{gatk}$	740	226	4588.77	58.93	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	740	4499	222.00	59.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	gatk	741	227	4498.77	58.91	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	$\operatorname{sam}$	741	4503	222.00	59.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	gatk	742	227	4566.77	58.91	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	742	4509	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	743	227	4633.77	58.91	0.00	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	743	4509	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	753	250	3750.77	58.29	0.78	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	753	4617	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	757	250	4258.77	58.29	34.57	Contaminants
Lmono	454	LGC	1	bwa	sam	757	4617	222.00	59.00		Contaminants
Lmono	454	LGC	1	TMAP	gatk	924	189	2175.77	43.35	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	926	189	2221.77	43.35	0.00	Contaminants
Lmono	454	$_{ m LGC}$	1	TMAP	gatk	928	189	2184.77	43.35	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	930	189	2184.77	43.35	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	953	250	4356.77	55.87	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	955	250	4334.77	55.87	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	957	250	4172.77	55.91	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	958	250	4210.77	55.91	0.00	Contaminants
Lmono	454	$_{ m LGC}$	1	TMAP	gatk	959	250	4251.77	55.91	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	961	250	4334.77	55.91	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	963	250	4217.77	55.91	0.00	Contaminants
Lmono	454	LGC	1	bwa	gatk	982	250	1899.77	60.00	687.88	Contaminants
Lmono	454	LGC	1	bwa	gatk	1047	250	3824.77	59.30	35.71	Contaminants
Lmono	454	LGC	1	bwa	sam	1047	8006	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1055	250	3067.77	59.16	2.72	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	sam	1055	8011	225.00	60.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	1072	250	3129.77	58.75	1.51	Contaminants
Lmono	454	LGC	1	bwa	sam	1072	8022	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	sam	1077	7975	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1192	250	5493.77	58.56	55.04	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	sam	1192	8008	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1201	250	4530.77	58.56	63.08	Contaminants
Lmono	454	LGC	1	bwa	sam	1201	8006	225.00	60.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	1208	250	3580.77	58.56	23.54	Contaminants
Lmono	454	LGC	1	bwa	sam	1208	8009	162.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1213	250	4330.77	58.56	64.05	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	sam	1213	8010	216.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1304	249	4878.77	59.84	69.84	Contaminants
Lmono	454	LGC	1	bwa	sam	1304	7998	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1307	250	5334.77	59.84	74.39	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	sam	1307	7999	225.00	60.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	1318	250	5106.77	59.78	91.22	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	$_{\mathrm{sam}}$	1318	8002	225.00	60.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	1321	250	4735.77	59.78	81.66	Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	$_{\mathrm{sam}}$	1321	8002	225.00	60.00		Contaminants
Lmono	454	$_{ m LGC}$	1	bwa	gatk	1329	250	4976.77	59.81	85.25	Contaminants

Lmono	454	LGC	1	bwa	sam	1329	8005	225.00	60.00		Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{gatk}$	1356	249	3918.77	59.73	50.60	Contaminants
Lmono	454	LGC	1	bwa	$\operatorname{sam}$	1356	8010	191.00	60.00		Contaminants
Lmono	454	LGC	2	bwa	gatk	315	250	3384.77	51.77	1.33	Contaminants
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	328	250	3396.77	51.77	1.33	Contaminants
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	346	250	149.77	59.51	37.09	End of read
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	347	250	413.77	59.51	51.04	End of read
Lmono	454	LGC	2	TMAP	gatk	347	250	130.77	91.19	40.53	End of read
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	555	144	69.77	57.88	2.20	Contaminants
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	587	144	83.77	57.88	2.17	Contaminants
Lmono	454	$_{\rm LGC}$	2	bwa	$\operatorname{gatk}$	677	145	122.77	59.80	2.17	Contaminants
Lmono	454	$_{\rm LGC}$	2	bwa	$\operatorname{gatk}$	700	145	119.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	gatk	703	145	119.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	gatk	712	145	105.77	59.80	2.17	Contaminants
Lmono	454	$_{\mathrm{LGC}}$	2	bwa	gatk	716	145	94.77	59.80	2.17	Contaminants
Lmono	454	$_{ m LGC}$	2	bwa	gatk	729	145	118.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	gatk	738	145	122.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	$_{ m gatk}$	740	145	122.77	59.80	2.17	Contaminants
Lmono	454	$_{ m LGC}$	2	bwa	$_{ m gatk}$	741	145	161.77	59.80	2.12	Contaminants
Lmono	454	$_{ m LGC}$	2	bwa	$_{ m gatk}$	742	145	119.77	59.80	2.17	Contaminants
Lmono	454	$_{ m LGC}$	2	bwa	gatk	743	145	119.77	59.80	2.17	Contaminants
Lmono	454	$_{ m LGC}$	$\overline{2}$	bwa	gatk	753	145	114.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	gatk	757	145	120.77	59.80	2.17	Contaminants
Lmono	454	LGC	2	bwa	gatk	963	122	701.29	60.00	0.00	Non-target region
Lmono	454	LGC	2	TMAP	gatk	963	21	82.31	74.82	0.00	Non-target region
Lmono	454	LGC	2	bwa	gatk	1047	250	3748.77	59.89	4.09	Contaminants
Lmono	454	LGC	2	bwa	gatk	1055	$\frac{250}{250}$	3571.77	59.89	7.83	Contaminants
Lmono	454	LGC	2	bwa	gatk	1072	$\frac{250}{250}$	2637.77	59.92	18.48	Contaminants
Lmono	454	LGC	$\frac{2}{2}$	bwa	gatk	1072	$\frac{230}{249}$	2161.77	59.92 $59.92$	18.66	Contaminants
Lmono	454	LGC	$\frac{2}{2}$	bwa	gatk	1192	$\frac{249}{250}$	4830.77	59.65	8.22	Contaminants
	454	LGC	$\frac{2}{2}$	bwa	_	1201	$\frac{250}{250}$	4741.77	59.65	5.67	Contaminants
Lmono	454	LGC	$\frac{2}{2}$		gatk	1201 $1208$	$\frac{250}{250}$	4741.77	59.65	8.23	Contaminants
Lmono				bwa	gatk						
Lmono	454	LGC	2	bwa	gatk	1213	250	4701.77	59.65	8.18	Contaminants
Lmono	454	LGC	2	bwa	gatk	1304	250	4652.77	60.00	6.46	Contaminants
Lmono	454	LGC	2	bwa	gatk	1307	250	4789.77	60.00	8.27	Contaminants
Lmono	454	LGC	2	bwa	gatk	1318	250	4710.77	60.00	8.42	Contaminants
Lmono	454	LGC	2	bwa	gatk	1321	250	4641.77	60.00	8.42	Contaminants
Lmono	454	LGC	2	bwa	gatk	1329	250	4555.77	60.00	10.65	Contaminants
Lmono	454	LGC	2	bwa	$\operatorname{gatk}$	1356	250	4453.77	60.00	8.63	Contaminants
Lmono	454	LGC	3	bwa	$\operatorname{gatk}$	346	250	102.77	60.00	32.12	End of read
Lmono	454	LGC	3	TMAP	gatk	346	250	294.77	91.54	40.01	End of read
Lmono	454	LGC	3	bwa	$\operatorname{gatk}$	347	250	255.77	60.00	42.21	End of read
Lmono	454	LGC	3	bwa	$\operatorname{gatk}$	370	250	67.77	60.00	8.83	End of read
Lmono	454	LGC	3	bwa	$\operatorname{gatk}$	963	111	302.48	60.00	0.00	Non-target region
Lmono	454	LGC	3	TMAP	$\operatorname{gatk}$	963	10	78.77	67.25	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	$\operatorname{gatk}$	330	250	5990.77	79.12	466.25	Strand bias
Lmono	454	NMIA	1	TMAP	$\operatorname{gatk}$	334	250	5973.77	79.12	514.24	Strand bias
Lmono	454	NMIA	1	TMAP	$\operatorname{gatk}$	335	250	5199.77	79.12	512.93	Strand bias
Lmono	454	NMIA	1	bwa	gatk	381	250	37.77	60.00	13.82	End of read
Lmono	454	NMIA	1	TMAP	gatk	533	249	1662.77	67.08	0.00	End of read
Lmono	454	NMIA	1	TMAP	$_{ m gatk}$	932	94	3555.77	75.44	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	$_{ m gatk}$	936	92	3557.77	75.89	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	954	250	8461.77	77.83	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	$_{ m gatk}$	957	250	8402.77	77.83	0.00	Non-target region
	-				0 .						G - 18 - 1

Lmono	454	NMIA	1	TMAP	gatk	961	250	8422.77	77.83	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	962	250	8332.77	77.83	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	963	250	8246.77	77.83	0.00	Non-target region
Lmono	ION	NIST	1	bwa	gatk	792	259	132.77	60.00	54.38	Strand bias
Lmono	ION	NIST	1	TMAP	gatk	792	275	323.77	85.79	69.53	Strand bias
Lmono	Sanger	LGC	1	bwa	$\operatorname{sam}$	390	81	25.50	60.00		End of read
Lmono	Sanger	LGC	1	bwa	$\operatorname{sam}$	1409	44	13.70	60.00		End of read
Lmono	Sanger	LGC	1	TMAP	$\operatorname{sam}$	1505	41	71.20	60.00		End of reference
Lmono	Sanger	LGC	1	TMAP	$\operatorname{sam}$	1506	41	71.20	60.00		End of reference
Lmono	Sanger	NIST	1	bwa	$\operatorname{sam}$	865	74	76.50	60.00		End of read
Lmono	Sanger	NIST	1	TMAP	$\operatorname{sam}$	865	68	77.50	59.00		End of read
Lmono	Sanger	NIST	1	bwa	gatk	867	67	264.77	60.00	0.00	End of read
Lmono	Sanger	NIST	1	bwa	$\operatorname{sam}$	867	67	10.40	60.00		End of read
Lmono	Sanger	NIST	1	TMAP	gatk	867	64	249.77	96.41	0.00	End of read
Lmono	Sanger	NIST	1	TMAP	$\operatorname{sam}$	867	64	12.30	59.00		End of read
Lmono	Sanger	NIST	1	TMAP	sam	1504	35	214.00	60.00		End of read

### Contaminants - BLAST results

BLAST reports for representative sequences of reads responsible for false positive variant calls in the LGC  $L.\ monocytogenes\ "454"$  rep 1 dataset.

#### BLASTN 2.2.29+

Reference: Zheng Zhang, Scott Schwartz, Lukas Wagner, and Webb Miller (2000), "A greedy algorithm for aligning DNA sequences", J Comput Biol 2000; 7(1-2):203-14.

RID: KH9SY3U8014

Database: Representative Chromosomes

2,857 sequences; 5,609,140,793 total letters

Query= Length=558

	Score	E
Sequences producing significant alignments:	(Bits)	Value
ref NC_000913.3  Escherichia coli str. K-12 substr. MG1655, c	979	0.0
ref NC_018658.1  Escherichia coli 0104:H4 str. 2011C-3493 chr	979	0.0
ref NC_017634.1  Escherichia coli 083:H1 str. NRG 857C chromo	979	0.0
ref NC_011751.1  Escherichia coli UMN026 chromosome, complete	979	0.0
ref NC_011750.1  Escherichia coli IAI39 chromosome, complete	979	0.0
ref NC_011740.1  Escherichia fergusonii ATCC 35469 chromosome	979	0.0
ref NC_007384.1  Shigella sonnei Ss046 chromosome, complete g	979	0.0
ref NC_002695.1  Escherichia coli 0157:H7 str. Sakai chromoso	979	0.0
ref NC_004337.2  Shigella flexneri 2a str. 301 chromosome, co	974	0.0
ref NC_007613.1  Shigella boydii Sb227 chromosome, complete g	974	0.0

#### ALIGNMENTS

>ref|NC\_000913.3| Escherichia coli str. K-12 substr. MG1655, complete genome

### Length=4641652

Features in this part of subject sequence: rRNA-16S ribosomal RNA of rrnH operon

Score = 979 bits (530), Expect = 0.0
Identities = 539/543 (99%), Gaps = 2/543 (0%)
Strand=Plus/Plus

Query	3	CCTGATGCAGCCATGCCGCGTGTATGAAGAAGGCTTACGGGTTGT-AAGTACGTTTCAGC 61	L
Sbjct	224155		24213
Query	62	GGGGAGGAAGGAATAAAGTTAATACCTTTGCTCATTGACGTTACCCGCAGAAGAAGCAC 12	21
Sbjct	224214		24273
Query	122	CGGCTAACTCCGTGCCAGCAGCCGCGGTAATACGGAGGGTGCAAGCGTTAATCGGAATTA 18	31
Sbjct	224274		24333
Query	182	CTGGGCGTAAAGCGCACGCAGGCGGTTTGTTAAGTCAGATGTGAAATCCCCGGGCTCAAC 24	11
Sbjct	224334		24393
Query	242	CTGGGAACTGCATCTGATACTGGCAAGCTTGAGTCTCGTAGAGGGGGGGTAGAATTCCAGG 30	)1
Sbjct	224394		24453
Query	302	TGTAGCGGTGAAATGCGTAGAGATCTGGAGGAATACCGGTGGCGAAGGCGGCCCCCTGGA 36	31
Sbjct	224454		24513
Query	362	CGAAGACTGACGCTCAGGTGCGAAAGCGTGGGGAGCAAACAGGATTAGATACCCTGGTAG 42	21
Sbjct	224514	CGAAGACTGACGCTCAGGTGCGAAAGCGTGGGGAGCAAACAGGATTAGATACCCTGGTAG 22	24573
Query	422	TCCACGCCGTAAACGATGTCGACTTGGAGGTTGTGCCCTTGAGGCGTGGCTTCCGGAGCT 48	31
Sbjct	224574	TCCACGCCGTAAACGATGTCGACTTGGAGGTTGTGCCCTTGAGGCGTGGCTTCCGGAGCT 22	24633
Query	482	AACGCGTTAAGTCGACCGCCTGGGGAGTACGGCCGCAAGGTTAAAACTCAAATGAATTGA 54	11
Sbjct	224634	AACGCGTTAAGTCGACCGCCTGGGGAGTACGGCCGCAAGGTTAAAACTCAAATGAATTGA 22	24693
Query	542	CGG 544	
Sbjct	224694	   CGG   224696	

Database: Representative Chromosomes
Posted date: Mar 21, 2014 12:17 AM
Number of letters in database: 5,609,140,793
Number of sequences in database: 2,857

```
Lambda
          K
                    Η
   1.33 0.621
                    1.12
Gapped
           K
Lambda
    1.28
           0.460
                     0.850
Matrix: blastn matrix:1 -2
Gap Penalties: Existence: 0, Extension: 0
Number of Sequences: 2857
Number of Hits to DB: 6177
Number of extensions: 6
Number of successful extensions: 6
Number of sequences better than 10: 1
Number of HSP's better than 10 without gapping: 0
Number of HSP's gapped: 3
Number of HSP's successfully gapped: 3
Length of query: 558
Length of database: 5609140793
Length adjustment: 30
Effective length of query: 528
Effective length of database: 5609055083
Effective search space: 2961581083824
Effective search space used: 2961581083824
A: 0
X1: 13 (25.0 bits)
X2: 32 (59.1 bits)
X3: 54 (99.7 bits)
S1: 13 (25.1 bits)
```

S2: 21 (39.9 bits)