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 the variants were potential false positives (Manuscript Table 2, tables 4 and 5). Consensus base quality statistics for biologically conserved positions (table 1).

Table 1: Biologically Conserved Position Base Qualities Characteristics of consensus based calls for conserved bases. Normalized quality values were obtained by dividing the quality values by the raw quality

score assigned by GATK foreach biologically conserved base position

	Di			D 0 1		3.51	3.5
Org	Plat	Lab	Rep	Raw Qual	Normalized	Min	Max
Ecoli	454	LGC	1	150854.23	2.99	1.26	3.01
Ecoli	454	LGC	2	71840.73	2.96	0.59	3.01
Ecoli	454	LGC	3	132329.73	2.98	1.17	3.01
Ecoli	454	NMIA	1	12612.23	2.69	0.38	3.02
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	19251.23	2.58	0.10	8.65
Lmono	454	LGC	2	151219.23	2.99	1.88	3.01
Lmono	454	LGC	3	133129.23	2.98	1.80	3.01
Lmono	454	NMIA	1	13444.73	2.61	1.00	8.65
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

A number of false positive variant calls were due to low sequencing coverage because the targeted sequencing strategy was responsible for false positive variant calls in six of the eight "454" datasets. For those six datasets, a variant was called at the last position in the gap between the two sequencing regions, bases 940 and 963 relative to reference sequences for *E. coli* and *L. monocytogenes*, respectively. A 40 bp region that was not part of the targeted sequencing region had significantly lower median coverage than the targeted region (2 X vs. 30,110 X, respectively) for all "454" datasets combined (Main Paper Figures 1 and 2).

A number of false postive variants were called due to contaminants. A low level of containating reads (150) present in the LGC L. monocytogenes rep 1 dataset. A BLAST analysis of a representative of these reads indicated that they were from E. coli (E value of 0.0), a well known contaminant of molecular biology reagents (section 4). A number of Escherichia coli strains E value of 0.0. in LGC Lmono rep 1 454 dataset. False positive variant calls were also attributable to the sequencing strategy and the variant calling algorithm. Resulting in a number of variants called due to strand bias. Stand bias was identified as the cause of the false positive variant call because greater than 99% of the reads were covering the variant bases were in the same direction. The strand bias was a product of the amplicon-based sequencing. For the E. coli dataset the variant was at the 3 end of the region 1 amplicon (Main Paper Figure 1) and the variant in the L. monocytogenes was as the 5 end of the region 2 amplicon (Main Paper Figure 2). As a result a majority of the reads covering the variants were in a single direction as the reads in the other direction were not long enough to cover the variant. For whole genome sequencing data, read direction biases can indicate a systematic error. The UnifiedGenotyper variant caller takes into consideration strand bias resulting in the false positive variant calls and reports strand bias using the Fisher exact test statistic (see Main Paper Table 2). A filtering step is commonly performed when calling SNPs that would have identified these as false positive variants due to the low number of reads with the variant base.

2 Biologically Variable Positions

To determine the ratio of bases at the biologically variant positions, a novel Bayesian analysis based on binomial sampling theory was developed. According to the binomial distribution, the observed base ratios, while precise (due to high coverage), differed significantly from all potential copy ratios. Subsequently given the observed base ratios a Bayesian approach was used to identify the most probable copy ratio out of the possible abundant base ratios assuming $E.\ coli$ and $L.\ monocytogenes$ have seven and six 16S gene copies respectively (figs. 1 and 2).

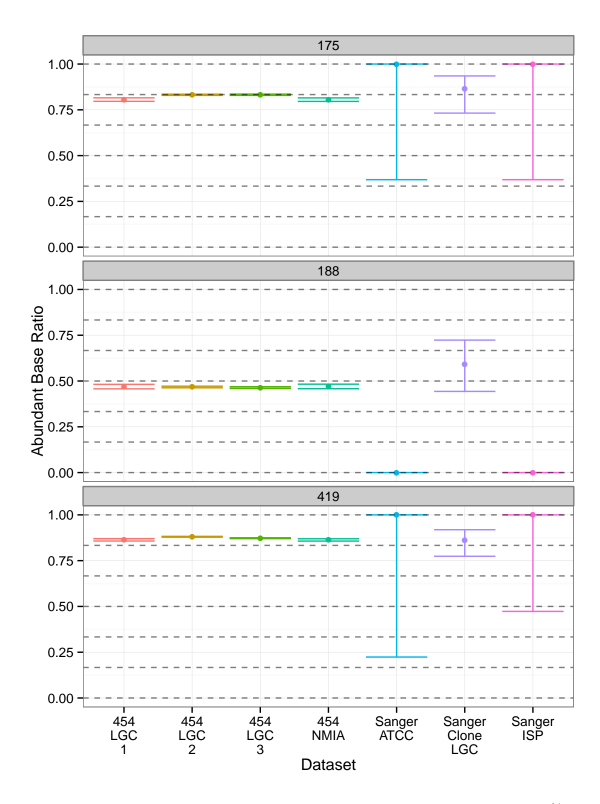


Figure 1: $L.\ monocytogenes$ biologically variable position base ratios. Error bars represent the 95 % posterior credibility interval estimated from a beta binomial distribution. Grey dashed lines indicate the potential base ratios assuming seven gene copies.

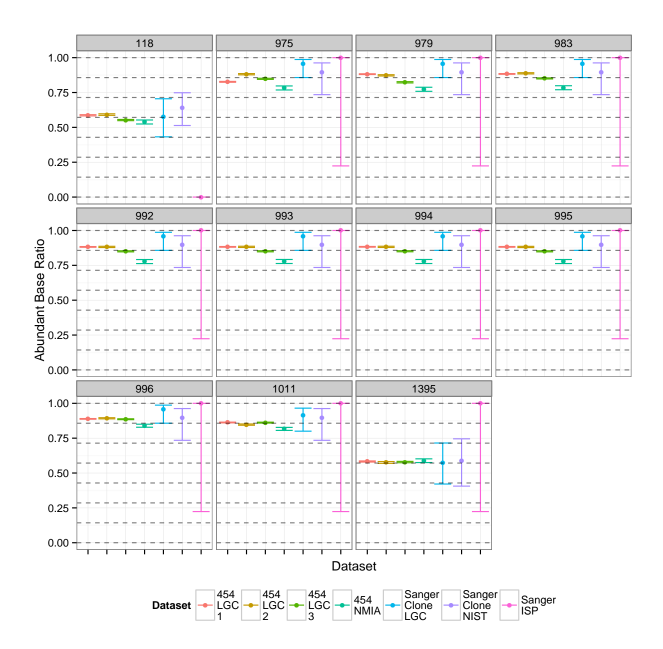


Figure 2: $E.\ coli$ biologically variable position base ratios. Error bars represent the 95 % posterior credibility interval estimated from a beta binomial distribution. Grey dashed lines indicate the potential base ratios assuming seven gene copies.

3 Likely sets of variant combinations

Most likely combination of variant strings for "454" and Sanger Clone library datasets table 2 and table 3).

Table 2: Estimated most likely set of variant combinations for E. coli

dataset	likelihood	chimera	ACCGATTGTA	ACCGATTGTG	GGTAGAATCA
Ecoli-454-LGC-1	0.04	353.03	3	3	1
Ecoli-454-LGC-2	0.03	258.60	3	3	1
Ecoli-454-LGC-3	0.08	33.30	3	3	1
Ecoli-454-NMIA-1	0.10	13.50	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	342.58	3	3	1

Table 3: Estimated most likely set of variant combinations for L. monocyotogenes

dataset	likelihood	chimera	GTA	GTG	TCA	TCG
Lmono-454-LGC-1	0.00	63.60	2	2	1	1
Lmono-454-LGC-2	0.01	658.52	2	2	1	1
Lmono-454-LGC-3	0.01	410.99	2	2	1	1
Lmono-454-NMIA-1	0.00	62.80	2	2	1	1
${\bf Lmono\text{-}LGC\text{-}Sanger\text{-}Clones.csv}$	0.00	5.13	2	2	1	1
Lmono-NIST-Sanger-Clones.csv	0.01	8.38	2	2	1	1
Consensus	0.01	1009.81	2	2	1	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.

Table 4: *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	
Ecoli	454	LGC	1	TMAP	gatk	324	250	423.77	88.92	61.22	Strand bias
Ecoli	454	LGC	1	bwa	gatk	325	250	308.77	60.00	53.48	
Ecoli	454	LGC	1	TMAP	gatk	325	250	300.77	88.92	53.96	
Ecoli	454	LGC	1	bwa	sam	396	2551	81.00	60.00		End of read
Ecoli	454	LGC	1	TMAP	sam	396	3013	38.00	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	19	215.77	85.03	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	124.77	69.56	0.00	
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	
Ecoli	454	LGC	3	TMAP	gatk	324	250	724.77	89.12	65.67	Strand bias
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	483.77	89.12	39.80	
Ecoli	454	LGC	3	bwa	gatk	348	250	741.77	59.92	11.62	End of read
Ecoli	454	LGC	3	bwa	sam	417	1032	22.00	60.00		Homopolymer
Ecoli	454	LGC	3	TMAP	sam	417	1020	33.00	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	233.29	85.63	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	313	250	5361.77	79.24	430.34	Strand bias
Ecoli	454	NMIA	1	TMAP	gatk	508	250	1204.77	83.50	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	509	250	1251.77	83.50	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	510	250	1345.77	83.50	0.00	End of read
Ecoli	454	NMIA	1	TMAP	gatk	514	250	1194.77	83.50	0.00	End of read
Ecoli	454	NMIA	1	TMAP	sam	514	6337	34.00	59.00		End of read
Ecoli	454	NMIA	1	TMAP	gatk	901	208	8102.77	84.35	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	904	208	8062.77	84.35	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	934	250	9479.77	75.40	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	935	250	9479.77	75.40	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	938	250	9312.77	75.35	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	sam	938	2746	9.54	56.00		Non-target region
Ecoli	454	NMIA	1	TMAP	gatk	939	250	9298.77	75.35	0.00	Non-target region
Ecoli	454	NMIA	1	TMAP	sam	939	2746	15.20	60.00		Non-target region
Ecoli	454	NMIA	1	TMAP	sam	941	2746	9.52	55.00		Non-target region
Ecoli	Sanger	NIST	1	TMAP	sam	1463	29	139.00	60.00		End of reference
Ecoli	Sanger	NIST	1	TMAP	sam	1464	29	214.00	60.00		End of reference

Table 5: *L. monocytogenes* Positions Pipeline Comparison Characteristics of variant calls for different bioinformatic pipelines.

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 bwa gatk 334 250 4678.77 70.65 302.04 Strand bias Lmono 454 LGC 1 bwa gatk 354 250 62.77 57.95 38.10 Lmono 454 LGC 1 bwa gatk 508 250 1386.77 51.13 7.03 Contaminar Lmono 454 LGC 1 bwa gatk 533 166 1407.77 46.04 0.00 Contaminar Lmono 454 LGC 1 bwa sam 533 250 2234.77 51.99 0.00 Contaminar Lmono 454 <t< th=""><th>ts ts ts ts ts ts ts ts</th></t<>	ts ts ts ts ts ts ts ts
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Lmono 454 LGC 1 bwa gatk 555 167 2077.77 46.14 0.00 Contaminar	$^{\mathrm{ts}}$
Lmono 454 LGC 1 bwa sam 555 1764 222.00 37.00 Contaminar	$^{\mathrm{ts}}$
Lmono 454 LGC 1 bwa gatk 559 167 2201.77 46.14 0.00 Contaminar	$^{\mathrm{ts}}$
Lmono 454 LGC 1 bwa sam 559 1763 222.00 37.00 Contaminar	ts
Lmono 454 LGC 1 bwa gatk 574 168 2288.77 46.03 0.00 Contaminar	ts
Lmono 454 LGC 1 bwa sam 574 1765 189.00 37.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 585 168 1737.77 46.03 0.00 Contaminar	
Lmono 454 LGC 1 bwa sam 585 1736 213.00 37.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 587 168 1980.77 46.03 0.00 Contaminar	
Lmono 454 LGC 1 bwa sam 587 1741 212.00 37.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 595 168 2352.77 46.03 0.00 Contaminar	ts
Lmono 454 LGC 1 bwa sam 595 1741 188.00 38.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 677 250 5127.77 58.81 4.08 Contaminar	ts
Lmono 454 LGC 1 bwa sam 677 4525 222.00 58.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 700 249 5372.77 56.79 14.63 Contaminar	
Lmono 454 LGC 1 bwa sam 700 4604 222.00 58.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 703 249 4820.77 56.79 4.37 Contaminar	
Lmono 454 LGC 1 bwa sam 703 4604 222.00 58.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 712 249 4896.77 56.79 13.22 Contaminar	
Lmono 454 LGC 1 bwa sam 712 4602 222.00 58.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 716 249 4009.77 56.79 0.00 Contaminar	
Lmono 454 LGC 1 bwa sam 716 4602 222.00 59.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 729 226 3856.77 58.93 0.72 Contaminar	
Lmono 454 LGC 1 bwa sam 729 4510 222.00 59.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 731 226 2873.77 58.93 0.00 Contaminar	
Lmono 454 LGC 1 bwa sam 731 4487 201.00 59.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 733 226 3028.77 58.93 0.00 Contaminar	
Lmono 454 LGC 1 bwa sam 733 4511 182.00 60.00 Contaminar	
Lmono 454 LGC 1 bwa gatk 738 226 4461.77 58.93 0.00 Contaminar	ts
Lmono 454 LGC 1 bwa sam 738 4493 222.00 59.00 Contaminar	

Lmono	454	$_{ m LGC}$	1	bwa	gatk	740	226	4588.77	58.93	0.00	Contaminants
Lmono	454	LGC	1	bwa	sam	740	4499	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	741	227	4498.77	58.91	0.00	Contaminants
Lmono	454	LGC	1	bwa	sam	741	4503	222.00	59.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	742	227	4566.77	58.91	0.00	Contaminants
Lmono	454	LGC	1	bwa	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	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	$_{\rm LGC}$	1	bwa	sam	753	4617	222.00	59.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	gatk	757	250	4258.77	58.29	34.57	Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	sam	757	4617	222.00	59.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	924	187	2152.77	44.40	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	924	616	6.19	11.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	926	187	2204.77	44.40	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	928	187	2122.77	44.40	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	930	187	2122.77	44.40	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	932	3	52.74	72.68	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	937	2	56.74	88.54	0.00	Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	gatk	939	2	56.74	88.54	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	952	250	630.77	53.23	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	953	250	4072.29	52.90	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	954	250	601.77	52.90	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	955	250	3489.77	52.90	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	957	250	3909.29	53.17	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	958	250	4719.77	53.17	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	959	250	3371.77	53.17	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	961	250	3413.77	53.17	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	962	250	640.77	53.17	0.00	Contaminants
Lmono	454	LGC	1	TMAP	gatk	963	250	3979.29	53.17	0.00	Contaminants
Lmono	454	LGC	1	bwa	gatk	982	250	1899.77	60.00	687.88	Contaminants
Lmono	454	LGC	1	TMAP	gatk	982	250	583.77	55.34	2.59	Contaminants
Lmono	454	LGC	1	TMAP	sam	1031	6732	3.01	42.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1035	6753	3.54	41.00		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	LGC	1	bwa	sam	1055	8011	225.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1055	8008	15.10	43.00		Contaminants
Lmono	454	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	TMAP	sam	1072	8013	27.00	44.00		Contaminants
Lmono	454	LGC	1	bwa	sam	1077	7975	225.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1077	7932	14.20	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1113	7985	14.20	43.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1114	8010	20.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1115	7987	19.10	43.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1116	8011	18.10	43.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1119	8007	9.52	43.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1145	8009	21.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1148	8011	16.10	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1150	8011	5.46	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1158	8011	16.10	44.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1192	250	5493.77	58.56	55.04	Contaminants

T	454	1.00	1	1		1100	0000	227 00	60.00		Ct
Lmono	$454 \\ 454$	$_{ m LGC}$	1 1	bwa TMAP	sam	$\frac{1192}{1192}$	8008 8011	$225.00 \\ 4.13$	60.00 43.00		Contaminants Contaminants
Lmono	454	LGC	1	bwa	sam	1192 1201	250	4.13 4530.77	58.56	63.08	Contaminants
Lmono	454	LGC	1	bwa bwa	gatk	1201 1201	8006	225.00	60.00	05.08	Contaminants
Lmono Lmono	454	LGC	1	TMAP	sam	1201 1201	8008	17.10	43.00		Contaminants
					sam					23.54	
Lmono	454	LGC	1	bwa	gatk	1208	250	3580.77	58.56	23.34	Contaminants
Lmono	454	LGC	1	bwa	sam	1208	8009	162.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1208	8012	18.10	44.00	04.05	Contaminants
Lmono	454	LGC	1	bwa	gatk	1213	250	4330.77	58.56	64.05	Contaminants
Lmono	454	LGC	1	bwa	sam	1213	8010	216.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1237	8002	24.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1238	8004	25.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1239	8004	4.77	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1248	8004	24.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1250	8003	16.10	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1257	8004	16.10	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1258	8004	14.20	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1266	8004	30.00	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1267	8001	12.30	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1268	8004	18.10	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1272	8002	12.30	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1273	8002	18.10	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1275	7999	7.80	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1277	8004	5.46	43.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1284	8004	12.30	43.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1285	8004	10.40	43.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1286	8004	3.01	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1287	8004	4.13	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1288	8000	4.13	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1291	8004	17.10	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1292	8004	17.10	44.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1296	8005	21.00	44.00		Contaminants
Lmono	454	LGC	1	bwa	gatk	1304	249	4878.77	59.84	69.84	Contaminants
Lmono	454	$_{ m LGC}$	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	LGC	1	bwa	sam	1307	7999	225.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1307	8005	3.01	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	gatk	1318	250	5106.77	59.78	91.22	Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	$_{ m sam}$	1318	8002	225.00	60.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	TMAP	sam	1318	8006	27.00	44.00		Contaminants
Lmono	454	$_{\rm LGC}$	1	bwa	gatk	1321	250	4735.77	59.78	81.66	Contaminants
Lmono	454	LGC	1	bwa	sam	1321	8002	225.00	60.00		Contaminants
Lmono	454	LGC	1	TMAP	sam	1321	8004	18.10	44.00		Contaminants
Lmono	454	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	gatk	1356	249	3918.77	59.73	50.60	Contaminants
Lmono	454	LGC	1	bwa	sam	1356	8010	191.00	60.00	00.00	Contaminants
Lmono	454	LGC	1	TMAP	sam	1356	8005	21.00	44.00		Contaminants
Lmono	454	LGC	2	bwa	gatk	315	250	3384.77	51.77	1.33	Contaminants
Lmono	454	LGC	2	bwa	gatk	328	250	3396.77	51.77	1.33	Contaminants
Lmono	454	LGC	2	bwa	gatk	$\frac{326}{346}$	$\frac{250}{250}$	149.77	59.51	37.09	Comaminants
Lmono	454	LGC	$\frac{2}{2}$	TMAP	gatk	346	$\frac{250}{250}$	277.77	91.87	46.11	
Lmono	454	LGC	$\frac{2}{2}$	bwa	gatk	347	$\frac{250}{250}$	413.77	59.51	51.04	
Lmono	454	LGC	$\frac{2}{2}$	TMAP	gatk	347	$\frac{250}{250}$	413.77 422.77	91.87	51.04 54.11	
LIHOHO	404	LGC	<u> </u>	INIAI	gath	941	200	444.11	91.01	04.11	

T	45.4	T 00	0	,	. 1		4.4.4	ao ==	*= 00	2.20	
Lmono	454	LGC	2	bwa	gatk	555	144	69.77	57.88	2.20	
Lmono	454	LGC	2	bwa	gatk	587	144	83.77	57.88	2.17	
Lmono	454	LGC	2	bwa	gatk	677	145	122.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	700	145	119.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	703	145	119.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	712	145	105.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	716	145	94.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	729	145	118.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	738	145	122.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	740	145	122.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	741	145	161.77	59.80	2.12	
Lmono	454	LGC	2	bwa	gatk	742	145	119.77	59.80	2.17	
Lmono	454	LGC	2	bwa	$_{ m gatk}$	743	145	119.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	753	145	114.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	757	145	120.77	59.80	2.17	
Lmono	454	LGC	2	bwa	gatk	963	122	701.29	60.00	0.00	Non-target region
Lmono	454	LGC	$\frac{2}{2}$	TMAP	gatk	963	21	82.31	74.82	0.00	Non-target region
Lmono	454	LGC	$\frac{2}{2}$	bwa	gatk	1047	250	3748.77	59.89	4.09	Contaminants
	454	LGC	$\frac{2}{2}$		_	1047 1055	$\frac{250}{250}$	3571.77	59.89	7.83	Contaminants
Lmono			$\frac{2}{2}$	bwa	gatk						
Lmono	454	LGC		bwa	gatk	1072	250	2637.77	59.92	18.48	Contaminants
Lmono	454	LGC	2	bwa	gatk	1077	249	2161.77	59.92	18.66	Contaminants
Lmono	454	LGC	2	bwa	gatk	1192	250	4830.77	59.65	8.22	Contaminants
Lmono	454	LGC	2	bwa	gatk	1201	250	4741.77	59.65	5.67	Contaminants
Lmono	454	LGC	2	bwa	gatk	1208	250	4714.77	59.65	8.23	Contaminants
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	$_{\rm LGC}$	2	bwa	gatk	1356	250	4453.77	60.00	8.63	Contaminants
Lmono	454	LGC	3	bwa	gatk	346	250	102.77	60.00	32.12	
Lmono	454	LGC	3	TMAP	gatk	346	250	285.77	91.99	40.01	
Lmono	454	LGC	3	bwa	gatk	347	250	255.77	60.00	42.21	
Lmono	454	LGC	3	bwa	gatk	370	250	67.77	60.00	8.83	
Lmono	454	LGC	3	bwa	gatk	963	111	302.48	60.00	0.00	Non-target region
Lmono	454	LGC	3	TMAP	gatk	963	10	78.77	67.25	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	330	250	5922.77	81.18	541.38	Strand bias
Lmono	454	NMIA	1	TMAP	gatk	334	$\frac{250}{250}$	5889.77	81.18	592.95	Strand bias
Lmono	454	NMIA	1	TMAP	gatk	335	$\frac{250}{250}$	5049.77	81.18	588.09	Strand bias Strand bias
					_					13.82	
Lmono	454	NMIA	1	bwa	gatk	381	250	37.77	60.00		End of read
Lmono	454	NMIA	1	TMAP	gatk	533	249	2139.77	69.73	0.00	End of read
Lmono	454	NMIA	1	TMAP	gatk	932	93	3553.77	74.69	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	936	91	3550.77	75.12	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	954	250	8664.77	78.57	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	957	250	8604.77	78.57	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	961	250	8684.77	78.58	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	962	250	8595.77	78.58	0.00	Non-target region
Lmono	454	NMIA	1	TMAP	gatk	963	250	8570.77	78.58	0.00	Non-target region
Lmono	Sanger	LGC	1	bwa	sam	390	81	25.50	60.00		End of read
Lmono	Sanger	LGC	1	bwa	sam	1409	44	13.70	60.00		End of read
Lmono	Sanger	LGC	1	TMAP	sam	1505	41	71.20	60.00		End of reference
Lmono	Sanger	LGC	1	TMAP	sam	1506	41	71.20	60.00		End of reference

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.