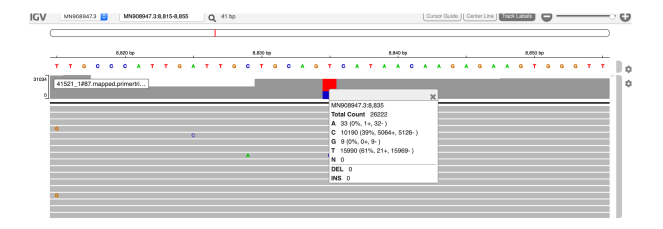
## Justification for template length filtering

## **Description of Issue**

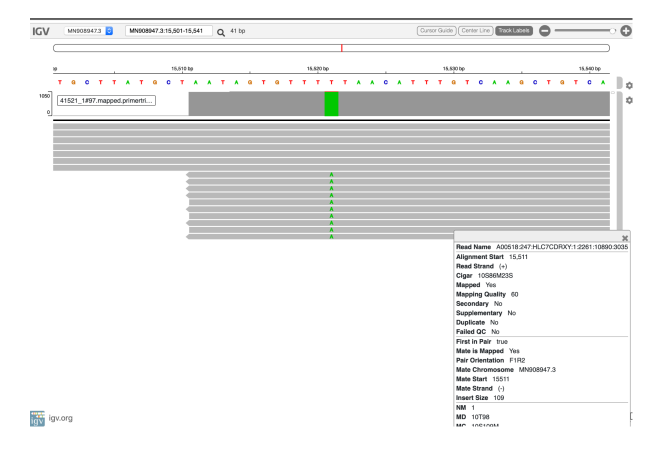
Unexpected SNPs in samples where primer panel V4 was used and this was linked to unexpected PCR products. In particular SNPs at positions 8835 (mutation orf1ab:V2857A, T -> C SNP) and 15521 (orf1ab:F5086Y)

Example 1 from IGV, sample 41521\_1#87 has a **T** -> **C** SNP at position 8835 in the variant calling file . This variant was called as it occured in 39% of reads, exceeding the 25% cutoff.

REGION	POS	REF	ALT	REF_DP	REF_RV	REF_QUA	λL	ALT_DP	ALT_RV	ALT_QUA	\L
ALT_FRE	Q	TOTAL_D	PVAL	PASS	GFF_FEA	TURE	REF_COI	OON	REF_AA	ALT_COD	ON
ALT_AA											
8835	T	С	15396	15375	35	9976	4945	36	0.393189	25372	0
TRUE	NA	NA	NA	NA	NA						

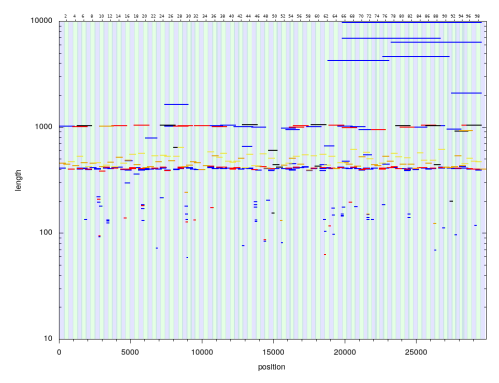


Example 2 from IGV , 41521\_1#97.mapped.primertrimmed.sorted.bam, T -> A SNP at 15521 in the short reads (mapping to reference position 15511 to 15596) is incorrect.



## Image showing presence of short length templates

## MILK-278E00D#87: Template sizes



## Proposed solution<sup>1</sup>

Filtering of short products less than 300 bp

samtools view -o \${sampleName}.filtered.mapped.bam -U \${sampleName}.bad.mapped.bam -e '!flag.munmap && (tlen > 300 || tlen < -300)' -F 2048

This should not remove genuine data as our minimum valid template is 385 bp

#### Results

A comparison of the consensus and variant calling pipeline results for run 41521 was done.

## Consensus file comparison

Comparing the 384 consensus files from run 41521 lane 1 before and after tlen filtering, 71 files were unchanged and 313 differed. The majority of the changes were an N in the filtered version<sup>2</sup>. Total positional changes across all samples were 4656 (1533 unique positions, 626 with >1 occurance). 158 of these were at reference position 8835. 284 at position 15521

#### Variant calling file comparison

Comparing the 384 ncovIlluminaCram\_ncovIllumina\_sequenceAnalysis\_callVariants tsv files from 41521 lane 1, 324/384 files had some changes in variants called.

There were 170 occurrences of T -> C at reference position 8835 in the original and none of these were in the filtered files. There were 301 occurrences of T -> A at reference position 15521, 3 of these were still present in the filtered file (low depth and/or FAIL)

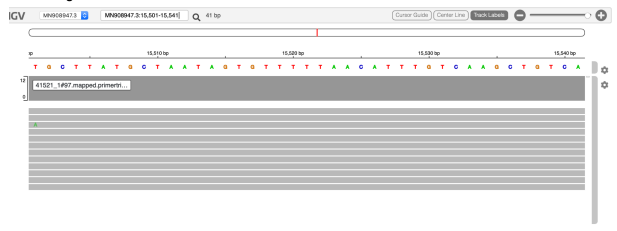
The majority of the other variant changes were due to SNPs removed after filtering. Of these, there were 37 across 32 samples. 33 were singleton SNPs. The 4 non-singletons were at position 71 (4 samples), 72 (11 samples), 14552 (2 samples) and 15451 (3 samples).

6 samples acquired a SNP after filtering. 5 of these were at position 28271 which was already widely observed in other samples. 1 from sample 41521\_1#383 was at 24825 and was low depth.

<sup>&</sup>lt;sup>1</sup> Proposed by Rob Davies, WSI

<sup>&</sup>lt;sup>2</sup> An exception was 41521\_1#106, which had a 35 bp deletion in the original file

Example from IGV post filtering, 41521\_1#97.mapped.primertrimmed.sorted.bam, reads containing the variant have been removed.



Example from IGV post filtering, 41521\_1#87.mapped.primertrimmed.sorted.filtered.bam, over 99% of remaining reads match the reference at position 8835

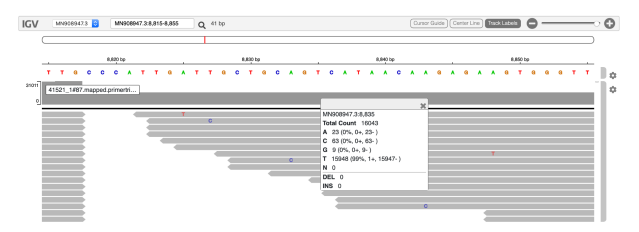
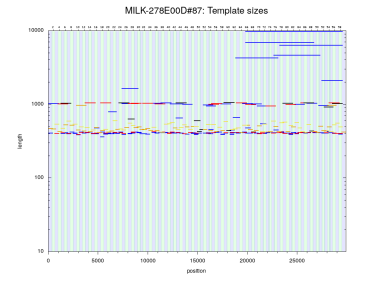


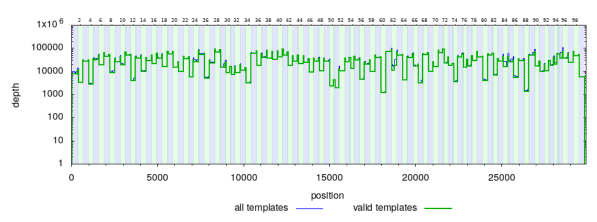
Image showing short templates having being removed.



## Valid templates appear unchanged

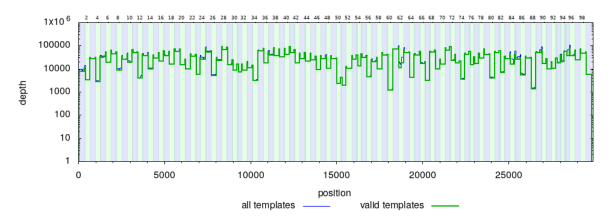
## Original





## Filtered

# MILK-278E00D#87: Template depth per base



# Artic pipeline duration

The time for the artic pipeline decreased for the filtered run. (Times in minutes)

	Min	Max	Mean	Median	Mode
Original:	1.07	34.73	14.04	14.48	16.28
Filtered:	1.02	30.83	10.48	10.83	9.69