# Analysis of viral barcodes in single-cell RNA-seq data.

#### David C. King and Fitz Fitzmeyer

### Synopsis

The following is an enumeration of distinct viral barcode sequences, taking into account read quality and considering the subsets defined by different samples (mosquito midguts), cells (cell barcode or whitelist), and UMIs (unique molecular identifiers). However, many distinct viral barcodes are attached to the same UMI with the same cell. This observation is unexpected and, at the time of writing, an anomalous observation that we consider unreliable until better understood. We can detect these anomalies by counting how many distinct viral sequences a UMI maps to within cells, and if this number is greater than 1, all reads for that midgut/cell/UMI combination are discarded.

#### Conclusion

In this dataset, the anomalous UMI/viral barcode sequences do not provide reliable evidence that more than one virus can infect a single cell. The phenomenon occurs at the highest quality standard (37), so cannot be attributed to base calling errors in the sequencing. Until the anomaly is explained, these observations must be filtered from the analysis, leaving few cases of deduplicated reads that indicate a distinct viral origin (ranging from 0 cases to 23 in different midgut samples). This analysis ignores some considerations. First, there may be evidence outside of the viral barcode that two distinct sequences share a UMI. If that is happening, the number of reliable counts would decrease further. Second, we did not evaluate whether this result is expected by UMI collision. However, an 11 base-pair UMI can have 4,194,304 unique sequences, far outnumbering the number of reads captured for a given cell. In conclusion, without being able to account for distinct sequences sharing a UMI in this dataset, we are unable to reliably observe viral diversity in a cell.

#### Data

There are 7 samples (West Nile virus infested mosquito midguts) represented by read pair data in fastq format. After searching for the viral barcode in the second read, the number of usable reads for this analysis is considerably less than the original file.

#### After selecting for reads with viral barcode

The script pysam\_analyze.py iterates over the matched read pair files and outputs the extracted barcodes, their quality string, and the minimum quality of each.

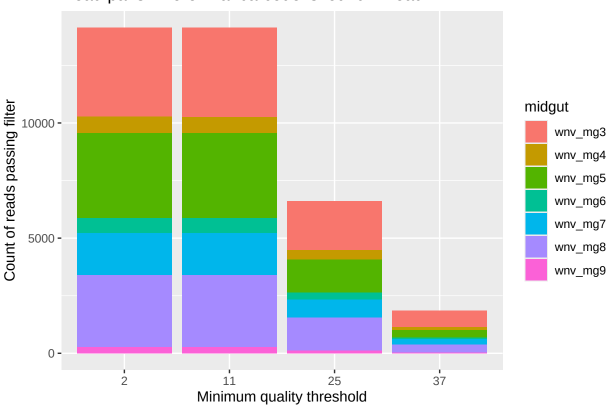
#### Summary of qualities per midgut

Table 1: Number of reads with the following lowest quality score

midgut	37	25	11	2
wnv_mg3	722	1413	1748	2
$wnv\_mg4$	137	271	292	1
$wnv\_mg5$	310	1117	2268	5
$wnv\_mg6$	84	219	335	2

midgut	37	25	11	2
wnv_mg7	220	564	1049	2
$wnv\_mg8$	352	1087	1677	5
$wnv\_mg9$	29	85	158	0

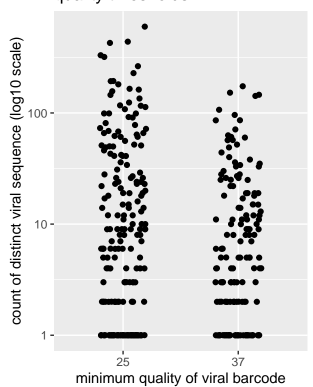
# Read pairs where viral barcode is found in read 2



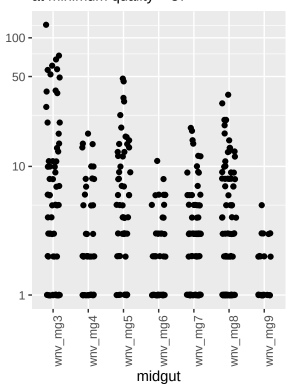
### Viral diversity

At the two top quality scores, 25 and 37, how many distinct viral barcode sequences do we see and how often do we see them? For each plot, a single point is a distinct viral barcode, its y value is the number of reads having that barcode, and it is distinct within the groups of the plot (x-axis).

# Distinct viral sequences at two quality thresholds



# Across midguts at minimum quality = 37



## Numbers for the above plots

Table 2: Breakdown of distinct viral counts by min. quality score

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	>= 25
																								59 33

Table 3: Breakdown of distinct viral counts by midgut at min. score of  $37\,$ 

																							>=
1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	25
mg3 17	5	6	2	5	3	2	2	1	3	3	0	1	1	1	0	0	1	0	0	0	2	0	12
mg4 16	12	3	0	3	1	1	3	0	2	0	0	0	1	2	0	0	1	0	0	0	0	0	0
mg5 11	4	6	5	3	3	2	2	2	1	0	2	2	1	2	1	2	0	0	1	0	0	0	5
mg6 22	7	7	2	2	5	0	1	0	0	1	0	0	0	0	0	0	0	0	0	0	0	0	0
mg7 34	9	19	3	6	4	1	0	3	1	0	2	0	0	1	1	0	0	1	1	0	0	0	0
mg8 27	11	8	4	4	1	3	7	3	2	1	1	3	2	0	1	0	1	0	0	1	0	3	2
mg9 16	5	4	0	1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0

#### Deduplication and the UMI

#### Background

cDNA libraries attempt to capture RNA molecules in a sample, such as mRNA transcripts produced during gene expression. To provide enough material for sequencing, the cDNA is usually amplified by PCR, boosting the "signal" of a molecular by increasing its representation in the isolated cDNA. The production of the RNA molecule is of biological origin. The duplication of the cDNA fragment is of technical origin. When the goal is to count the number of molecules produced by biological means, the technical duplication should be controlled for.

The current approach is to use a unique molecular identifier (UMI), a short oligonucleotide attached to the cDNA of a biologically produced molecule *before* the amplification step. Therefore, identical sequences which include the same UMI are assumed to come from a single molecule and are processed with software to collapse, or deduplicate, the duplicate sequences into a single observation.

In actual sequenced reads, there can be variation in the observed sequence due to sequencing errors or uncertainty in the base call. Therefore, some ambiguity must be taken into account when deduplicating reads. Most methods allow a small number of mismatches between UMIs if the attached sequence maps to the same place in the target genome. Therefore, with the aid of the alignment, UMIs and the read can have variation.

Given any two sequence reads, the following cases are treated this way: - Same UMI; same sequence: single molecule duplicated by PCR - Different UMI; same sequence: two distinct molecules generated biologically (e.g. two mRNAs transcribed from a gene) - Different UMI; different sequence: If UMIs are close (i.e. <= 1 mismatch) AND the attached sequences map to exactly the same place via alignment, the two reads may have been a PCR duplication. - Same UMI; different sequence: ??????????????????

#### Anomalous UMI-sequence combinations

Grouping by distinct combinations of UMI and payload will determine uniqueness.

Filter first by quality >= 37, then count instances after grouping by distinct UMIs. There is no deduplication, so "occurence" includes PCR duplicate read counts. No breakdown by yet midgut.

Distinct viral barcode sequences	s that share a UMI	occurrence
1		3277
2		102
3		30
4		19
5		13
6		10
7		8
8		3
9		5
10		5
>= 10		106

Table 4: How many different viral barcodes share a distinct UMI?

#### Break apart by midgut

#### Viral diversity after deduplicating

After filtering out the anomalous cases, deduplication can be performed by collapsing exact copies of UMI:viral barcode reads into a single row in the table.

# Count of distinct viral sequences by midgut AFTER 'dedup' Minimum quality = 37

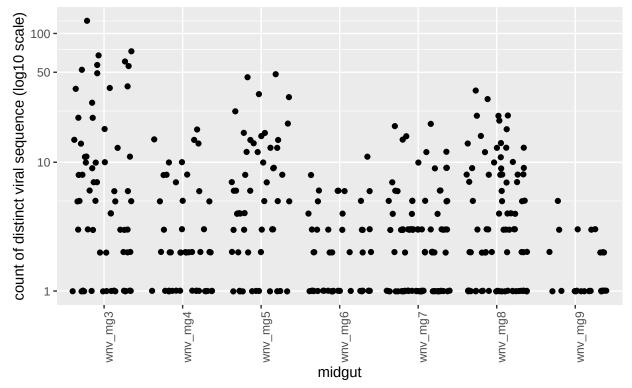


Table 5: data for figure above

	$wnv\_mg3$	$wnv\_mg4$	$wnv\_mg5$	$wnv\_mg6$	$wnv\_mg7$	$wnv\_mg8$	$wnv\_mg9$
1	17	16	11	22	34	27	16
2	5	12	4	7	9	11	5
3	6	3	6	7	19	8	4
4	2	0	5	2	3	4	0
5	5	3	3	2	6	4	1
6	3	1	3	5	4	1	0
7	2	1	2	0	1	3	0
8	2	3	2	1	0	7	0
9	1	0	2	0	3	3	0
>=10	24	6	17	1	7	17	0

## Viral diversity within cells

#### Break out by cell barcode

```
## # A tibble: 6 x 8
               midgut, cellbarcode [5]
## # Groups:
     midgut cellbarcode
                             umi
                                          viral_seq
                                                        distinct_viral_seque~1
##
     <chr>
             <chr>>
                              <chr>
                                          <chr>
                                                                          <int> <int>
## 1 wnv_mg3 AAACCTGCGCTAACC TGTACTGCCTT CTAACCGTAACT~
                                                                              1
                                                                                    1
## 2 wnv_mg3 AAACCTGCGCTAACC TGTACTGTCTT CTAACCGTAACT~
                                                                              1
                                                                                    1
## 3 wnv_mg3 AAACCTGCTTCTCGT ACTGTCTTCAC CTAACTGTAACC~
                                                                              1
                                                                                    1
## 4 wnv_mg3 AAACCTGGTGATAAA CTCGGCCCAGG CTTACCGTAACG~
                                                                                    1
```

```
## 5 wnv_mg3 AAACGGGAACCATGT AGGCATAGTGG CTTACCGTTACT~
                                                                                     1
## 6 wnv_mg3 AAACGGGGTCCATCC TTTCTTCATAG CTAACTGTAACC~
## # i abbreviated name: 1: distinct_viral_sequences
## # i 2 more variables: midgut_cellbarcode <chr>, umi_viral_seq <chr>
## .
                                7
##
      1
           2
                3
                     4
                           6
## 5146
          45
                5
                     1
                           1
##
##
      1
           2
                3
                     4
## 5108
          42
                     1
```

# Cells with more than one distinct virus no whitelisting

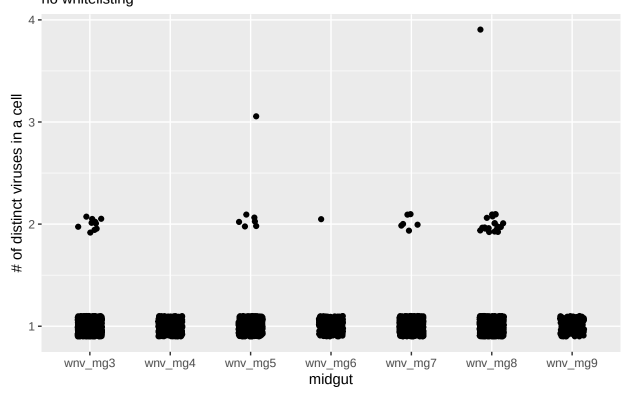


Table 6: Numbers for the plot above

	$wnv\_mg3$	$wnv\_mg4$	$wnv\_mg5$	$wnv\_mg6$	$wnv\_mg7$	$wnv\_mg8$	$wnv\_mg9$
1	1309	415	880	332	794	1194	184
2	10	0	6	1	6	19	0
3	0	0	1	0	0	0	0
4	0	0	0	0	0	1	0

```
## # A tibble: 6 x 8
## # Groups: midgut, cellbarcode_consensus_1 [5]
                                             viral_seq distinct_viral_seque~1
     midgut cellbarcode_consensus_1 umi
             <chr>>
##
     <chr>
                                     <chr>
                                             <chr>>
                                                                         <int> <int>
## 1 wnv_mg3 AAACCTGCGCTAACC
                                     TGTACT~ CTAACCGT~
                                                                             1
                                                                                   1
## 2 wnv_mg3 AAACCTGCGCTAACC
                                    TGTACT~ CTAACCGT~
                                                                             1
                                                                                   1
## 3 wnv_mg3 AAACCTGGTGATAAA
                                    CTCGGC~ CTTACCGT~
                                                                             1
                                                                                   1
```

```
## 4 wnv_mg3 AAACGGGAACCATGT
                                      AGGCAT~ CTTACCGT~
                                                                               1
                                                                                      1
                                                                               1
## 5 wnv_mg3 AAACGGGGTCCATCC
                                      TTTCTT~ CTAACTGT~
                                                                                      1
                                      CTACTG~ CTGACAGT~
                                                                               1
                                                                                      2
## 6 wnv_mg3 AAACGGGGTTCACTA
## # i abbreviated name: 1: distinct_viral_sequences
## # i 2 more variables: midgut_cellbarcode <chr>, umi_viral_seq <chr>
##
                                7
##
      1
           2
                3
                     4
                           6
## 4370
          43
                6
                      1
                           1
```

### Cells with more than one distinct virus

UMI anomalies removed; cell barcode whitelist (mismatch <= 1)

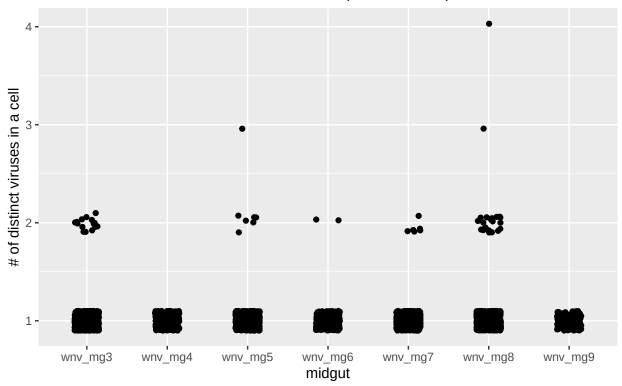


Table 7: Numbers for the plot above

	$wnv\_mg3$	$wnv\_mg4$	$wnv\_mg5$	$wnv\_mg6$	$wnv\_mg7$	wnv_mg8	wnv_mg9
1	1134	346	742	273	663	1003	151
2	15	0	6	2	6	21	0
3	0	0	1	0	0	1	0
4	0	0	0	0	0	1	0

#### Conclusion

In this dataset, the anomalous UMI/viral barcode sequences do not provide reliable evidence that more than one virus can infect a single cell. The phenomenon occurs at the highest quality standard (37), so cannot be attributed to base calling errors in the sequencing. Until the anomaly is explained, these observations must be filtered from the analysis, leaving few cases of deduplicated reads that indicate a distinct viral origin (ranging from 0 cases to 23 in different midgut samples). This analysis ignores some considerations. First, there may

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