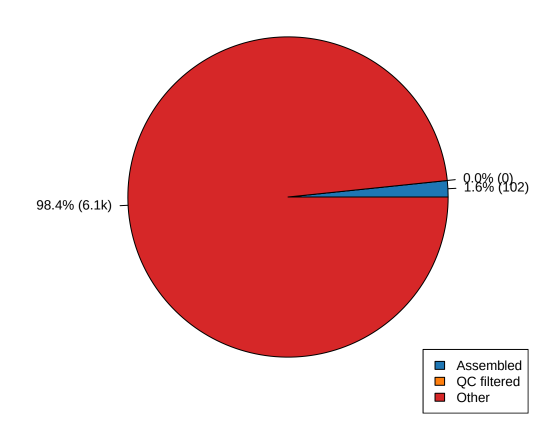
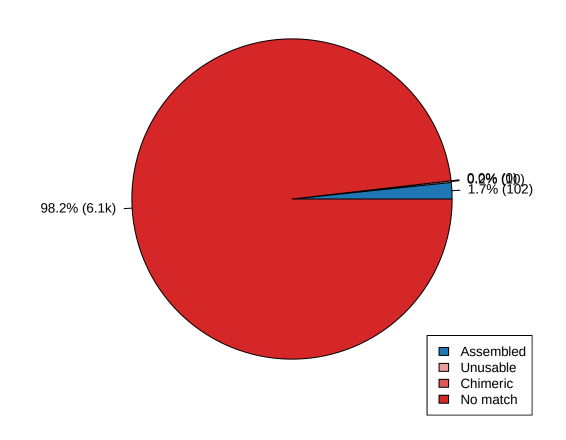
### 1. Percentages of total reads

### 2. Percentages of all read patterns passing QC





## 3. Percentages of assembled reads

# 33.3% (34) A\_MP<sub>1</sub> 9.8% (10) A\_HA\_H1 5.9% (6) A\_PB2 2.9% (3) A\_NP 26.5% (27) A\_NS

### **SAMPLE** "252232567\_N"

#### READ PROPORTIONS.

- 1. Percentages of total read counts
  - ASSEMBLED: influenza reads in final assemblies.
  - QC FILTERED: didn't pass length/median quality thresholds.
  - OTHER: non-flu and contaminant/poor flu signal.
- 2. Percentages of all read patterns passing QC process
  - Patterns are clustered or non-redundant reads.
  - ASSEMBLED: excellent influenza read patterns.
  - UNUSABLE: poor or contaminant flu patterns.
  - CHIMERIC: flu patterns matching both strands.
  - NO MATCH: non-flu read patterns.
- 3. Percentages of assembled read counts
- Shows the proportion of gene segments to the genome.
- Paired-end reads have been merged into a single count unless not applicable: single-end reads have been used.