

1 Extreme heterogeneity of influenza 2 virus infection in single cells

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7

8 **Abstract** Viral infection can dramatically alter a cell's transcriptome. However, these changes
9 have mostly been studied by bulk measurements on many cells. Here we use single-cell mRNA
10 sequencing to examine the transcriptional consequences of influenza virus infection. We find
11 extremely wide cell-to-cell variation in the productivity of viral transcription – viral transcripts
12 comprise less than a percent of total mRNA in many infected cells, but a few cells derive over half
13 their mRNA from virus. Some infected cells fail to express at least one viral gene, but this gene
14 absence only partially explains variation in viral transcriptional load. Despite variation in viral load,
15 the relative abundances of viral mRNAs are fairly consistent across infected cells. Activation of
16 innate immune pathways is rare, but some cellular genes co-vary in abundance with the amount of
17 viral mRNA. Overall, our results highlight the complexity of viral infection at the level of single cells.

18

19 Introduction

20 Viruses can cause massive and rapid changes in a cell's transcriptome as they churn out viral mRNAs
21 and hijack cellular machinery. For instance, cells infected with influenza virus at high multiplicity
22 of infection (MOI) express an average of 50,000 to 100,000 viral mRNAs per cell, corresponding
23 to 5 to 25% of all cellular mRNA (Hatada et al., 1989). Infection can also trigger innate-immune
24 sensors that induce the expression of cellular anti-viral genes (Killip et al., 2015; Iwasaki and Pillai,
25 2014; Crotta et al., 2013). This anti-viral response is another prominent transcriptional signature of
26 high-MOI influenza virus infection in bulk cells (Geiss et al., 2002).

27 However, initiation of an actual influenza infection typically involves just a few virions infecting
28 a few cells (Varble et al., 2014; Poon et al., 2016; Leonard et al., 2017; McCrone et al., 2017). The
29 dynamics of viral infection in these individual cells may not mirror bulk measurements made
30 on many cells infected at high MOI. Over 70 years ago, Max Delbrück showed that there was a
31 ~100-fold range in the number of progeny virions produced per cell by clonal bacteria infected
32 with clonal bacteriophage (Delbrück, 1945). Subsequent work has shown similar heterogeneity
33 during infection with other viruses (Zhu et al., 2009; Schulte and Andino, 2014; Combe et al., 2015;
34 Akpinar et al., 2016), including influenza virus (Heldt et al., 2015).

35 In the case of influenza virus infection, targeted measurements of specific proteins or RNAs
36 have shed light on some factors that contribute to cell-to-cell heterogeneity. The influenza virus
37 genome consists of eight negative-sense RNA segments, and many infected cells fail to express
38 one more of these RNAs (Heldt et al., 2015; Dou et al., 2017) or their encoded proteins (Brooke
39 et al., 2013). In addition, activation of innate-immune responses is inherently stochastic (Shalek
40 et al., 2013, 2014; Bhushal et al., 2017; Hagai et al., 2017), and only some influenza-infected cells
41 express anti-viral interferon genes (Perez-Cidoncha et al., 2014; Killip et al., 2017). However, the
42 extent of cell-to-cell variation in these and other host and viral factors remains unclear, as does the

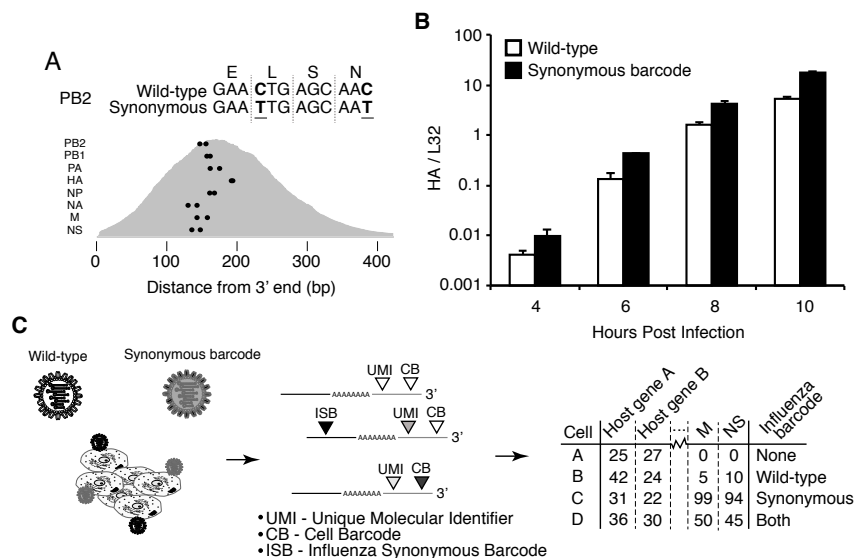


Figure 1. Experimental design. **(A)** We engineered a virus that carried two synonymous mutations near the 3' end of each mRNA. At top are the mutations for PB2. At bottom are locations of the synonymous mutations relative to the typical distribution of read depth for our 3'-end sequencing. **(B)** The wild-type and synonymously barcoded viruses transcribe their genes with similar kinetics. The abundance of the viral hemagglutinin (HA) transcript relative to the cellular housekeeping gene L32 was assessed by qPCR in A549 cells infected at an MOI of 0.5 (as determined on MDCK-SIAT1 cells). Error bars \pm S.D., n=3. **(C)** For the single-cell mRNA sequencing, A549 cells were infected with an equal mixture of wild-type and synonymously barcoded virus. Immediately prior to collection, cells were physically separated into droplets and cDNA libraries were generated containing the indicated barcodes. The libraries were deep sequenced, and the data processed to create a matrix that gives the number of molecules of each transcript observed in each cell. Infected cells were further annotated by whether their viral mRNAs derived from wild-type virus, synonymously barcoded virus, or both.

Figure 1-source data 1. Sequences of wild-type and barcoded viruses are in [viralsequences.fasta](#).

43 association among them in individual infected cells.

44 Here we use single-cell mRNA sequencing to quantify the levels of all cellular and viral mRNAs
45 in cells infected with influenza virus at low MOI. We find extremely large variation in the amount
46 of viral mRNA expressed in individual cells. Both co-infection and activation of innate-immune
47 pathways are rare in our low-MOI infections, and do not appear to be the major drivers of cell-
48 to-cell heterogeneity in viral transcriptional load. Individual infected cells often fail to express
49 specific viral genes, and such gene absence explains some but certainly not all of the cell-to-cell
50 heterogeneity. A variety of cellular genes, including ones involved in the oxidative-stress response,
51 co-vary with viral transcriptional load. Overall, our work demonstrates remarkable heterogeneity in
52 the transcriptional outcome of influenza virus infection among nominally identical cells infected
53 with a relatively pure population of virions.

54 Results

55 Strategy to measure mRNA in single virus-infected cells.

56 We performed single-cell mRNA sequencing using a droplet-based system that physically isolates
57 individual cells prior to reverse transcription ([Zheng et al., 2017; Macosko et al., 2015; Klein et al., 2015](#)).
58 Each droplet contains primers with a unique *cell barcode* that tags all mRNAs from that
59 droplet during reverse-transcription. Each primer also contains a *unique molecular identifier (UMI)*
60 that is appended to each mRNA molecule during reverse transcription. The 3' ends of the mRNAs are
61 sequenced and mapped to the human and influenza virus transcriptomes to determine transcript
62 identities. This information is combined with that provided by the UMIs and cell barcodes to

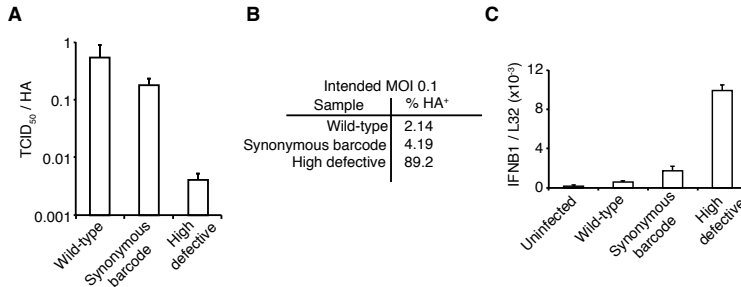


Figure 2. The viral stocks in our experiments are relatively pure of defective particles. **(A)** Our viral stocks have a higher ratio of infectious particles to HA virion RNA compared to a high-defective stock propagated at high MOI. HA viral RNA was quantified by qPCR on virions. Error bars \pm S.D., n=6 (qPCR replicates). **(B)** Our viral stocks have a higher ratio of infectious particles to particles capable of expressing HA protein. A549 cells were infected at an MOI of 0.1, and the percentage of cells expressing HA protein at 9 hours post-infection was quantified by antibody staining and flow cytometry. **(C)** Our viral stocks are less immunostimulatory than virus propagated at high MOI when used at the same number of infectious units as calculated by TCID₅₀. Note that this fact does not necessarily imply that they are more immunostimulatory per virion, as the high-MOI stocks also have more virions per infectious unit as shown in the first two panels. Measurements of *IFNB1* transcript by qPCR normalized to the housekeeping gene L32 in A549 cells at 10 hours post infection at an MOI of 0.5. Error bars \pm S.D., n=3. Note that MOIs were calculated by TCID₅₀ on MDCK-SIAT1 cells, whereas the experiments in this figure involved infection of A549 cells.

Figure 2-Figure supplement 1. Full flow cytometry data for panel B .

63 quantify the number of molecules of each mRNA species that have been captured for each cell.
 64 Infected cells will express viral as well as cellular mRNAs – however the cell barcodes and UMIs
 65 cannot distinguish whether a cell was initially infected by one or multiple viral particles. We therefore
 66 engineered an influenza virus (strain A/WSN/1933) that additionally carried *viral barcodes* consisting
 67 of synonymous mutations near the 3' end of each transcript (Figure 1A). Critically, these synonymous
 68 mutations did not greatly impact viral growth kinetics (Figure 1B). We infected A549 human lung
 69 carcinoma cells with an equal mix of the wild-type and synonymously barcoded viruses. Cells
 70 infected by a single virion will exclusively express mRNAs from either wild-type or synonymously
 71 barcoded virus, whereas cells that are co-infected with multiple virions will often express mRNAs
 72 from both the wild-type and synonymously barcoded viruses (Figure 1C).

73 We took care to generate stocks of virus that were relatively “pure” of defective particles. Stocks
 74 of viruses typically contain an array of biologically active viral particles, some of which are defective
 75 for replication owing to mutations or deletions in essential viral genes (*von Magnus, 1954; Huang*
76 et al., 1970; Brooke, 2014; Fonville et al., 2015; Lauring and Andino, 2010; Dimmock et al., 2014;
77 Saira et al., 2013). These defective particles become prevalent when a virus is grown at high MOI,
 78 where complementation permits the growth of otherwise deleterious genotypes. To minimize the
 79 levels of defective particles, we propagated our viral stocks at low MOI for a relatively brief period of
 80 time (*Xue et al., 2016*). We validated that our stocks exhibited greater purity of infectious particles
 81 than a stock propagated at high MOI by verifying that they had a higher ratio of infectious particles
 82 to virion RNA (Figure 2A) and to particles capable of inducing expression of a single viral protein
 83 (Figure 2B). In addition, viral stocks with many defective particles are more immunostimulatory
 84 per infectious unit (e.g., TCID₅₀) than low-defective stocks (*Tapia et al., 2013; Lopez, 2014*), in part
 85 simply because there are more physical virions per infectious unit (Figure 2A,B). We confirmed that
 86 our viral stocks induced less interferon per infectious unit than a stock propagated at higher MOI
 87 (Figure 2C).

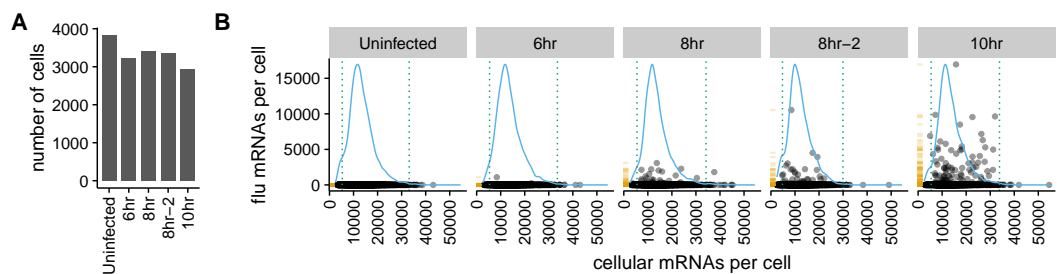


Figure 3. There is a very wide distribution in the amount of viral mRNA per cell. **(A)** Number of cells sequenced for each sample. **(B)** The number of cellular and viral mRNAs detected for each cell is plotted as a point. The blue lines show the overall distribution of the number of cellular mRNAs per cell. The orange rug plot at the left of each panel shows the distribution of the number of viral mRNAs per cell. Cells outside the dotted green lines were considered outliers with suspiciously low or high amounts of cellular mRNA (possibly derived from two cells per droplet), and were excluded from all subsequent analyses. Figure 3-Figure supplement 1 shows the exact distributions of the fraction of viral mRNA per cell.

Figure 3-Figure supplement 1. Cumulative fraction plot of proportion of total mRNA from virus.

88 Single cells show an extremely wide range of expression of viral mRNA.

89 We infected A549 cells at low MOI with a mixture of the wild-type and synonymously barcoded
90 viruses, and collected cells for sequencing at 6, 8, and 10 hours post-infection, performing two
91 slightly different variants of the experiment for the 8-hour timepoint. For most of the samples, we
92 replaced the infection inoculum with fresh media at one-hour post-infection, thereby ensuring that
93 most infection was initiated during a narrow time window. However, for the second 8-hour sample
94 (which we denote as "8hr-2" in the figures), we did *not* perform this media change and instead left
95 the cells in the original infection inoculum. The rationale for including a sample without a media
96 change was to determine the importance of synchronicity of the timing of infection as discussed
97 later in this subsection.

98 We recovered between 3,000 and 4,000 cells for each sample (Figure 3A). As expected for a
99 low-MOI infection, most cells expressed little or no viral mRNA (Figure 3B and Figure 3-Figure sup-
100 plement 1). Also as expected, the amount of viral mRNA per cell among infected cells increased
101 over time (Figure 3B, Figure 3-Figure supplement 1). But what was most notable was how widely the
102 number of viral mRNA molecules varied among infected cells. While the fraction of mRNA derived
103 from virus was <0.1% for most cells, viral mRNA constituted half the transcriptome in a few cells at
104 8 and 10 hours (Figure 3B, Figure 3-Figure supplement 1).

105 A complicating factor is that uninfected cells could have small amounts of viral mRNA due
106 to leakage of transcripts from lysed cells. It is therefore important to establish a threshold for
107 identifying truly infected cells. We can do this by taking advantage of the fact that roughly half the
108 infecting virions bear synonymous barcodes. Reads derived from lysed cells will be drawn from
109 both wild-type and synonymously barcoded viral transcripts. However, most cells are infected by at
110 most one virion, and so the reads from truly infected cells will usually derive almost entirely from
111 one of the two viral variants. Figure 4A shows the fraction of viral reads in individual cells from each
112 viral variant, and Figure 4B indicates the fraction of viral reads from the most abundant variant in
113 that cell. Most cells with large amounts of viral mRNA have viral transcripts exclusively derived from
114 one viral variant – indicating non-random partitioning as expected from viral infection. However,
115 cells with a small amount of viral mRNA often have viral transcripts from both variants, as expected
116 from the random partitioning associated with simple mRNA leakage. Finally, a few cells with large
117 amounts of viral mRNA have viral transcripts from both variants, likely reflecting co-infection.

118 We determined the threshold amount of viral mRNA per cell for each sample at which the
119 barcode partitioning clearly resulted from infection rather than leakage (Figure 4C and Figure 4-
120 Figure supplement 2), and used these thresholds to annotate cells that we were confident were
121 truly infected. We also annotated as co-infected cells above this threshold that had mRNA from

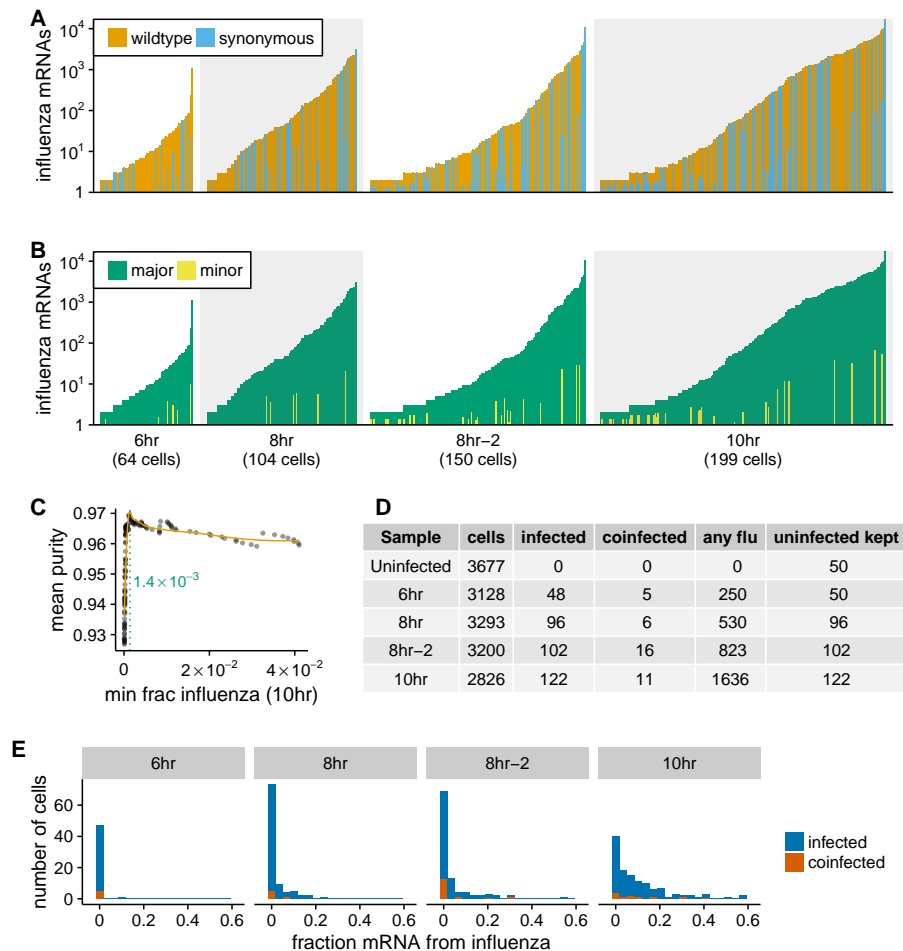


Figure 4. Synonymous barcodes on the viral mRNAs distinguish true infections from cells that contain viral mRNAs derived from leakage of lysed cells. **(A)** Cells with at least two viral mRNAs for which the barcode could be called, arranged in order of increasing influenza transcript counts. Bar heights denote the number viral mRNAs on a \log_{10} scale, bar coloring is linearly proportional to the fractions of viral mRNAs derived from wild-type and synonymously barcoded virus. **(B)** Same as (A), but each bar is colored according to the relative fraction of the more common (major) and less common (minor) virus variant. At low levels of viral mRNA there is often a roughly equal mix, suggesting contamination with viral mRNAs leaked from lysed cells. At higher levels of viral mRNA, cells generally have only one viral variant, suggesting infection initiated by a single virion. A few cells are also obviously co-infected with both viral variants. **(C)** We determined a threshold for calling “true” infections by finding the amount of viral mRNA per cell at which the viral barcode purity no longer increases with more viral mRNA. The purity is the fraction of all viral mRNA in a cell derived from the most abundant viral barcode in that cell. We fit a curve (orange line) to the mean purity of all cells with more than the indicated amount of viral mRNA, and drew the cutoff (dotted green line) at the point where this curve stopped increasing with the fraction of total mRNA derived from virus. This plot illustrates the process for the 10-hour sample, see Figure 4-Figure supplement 2 for similar plots for other samples. See the Methods for details. **(D)** The number of cells identified as infected and co-infected for each sample, as well as the number of cells with any viral read. For all subsequent analyses, we subsampled the number of uninfected cells per sample to the greater of 50 or the number of infected cells. **(E)** Distribution of the fraction of mRNA per cell derived from virus for both infected and co-infected cells. Figure 4-Figure supplement 3 shows these same data in a cumulative fraction plots and calculates Gini coefficients to quantify the heterogeneity in viral mRNA load.

Figure 4-Figure supplement 1. Number of viral barcodes called.

Figure 4-Figure supplement 2. Thresholds for calling infected cells.

Figure 4-Figure supplement 3. Cumulative distributions of viral mRNA per cell and Gini coefficients.

Figure 4-Figure supplement 4. Synchronization of infection does not greatly affect heterogeneity.

Figure 4-Figure supplement 5. Effects of infectious dose or coinfection state.

122 both viral variants. Figure 4D shows the number of cells annotated as infected and co-infected for
 123 each sample – these cells are just a small fraction of the number of cells with any viral read. These
 124 annotation thresholds are conservative, and may miss some true low-level infections. However, it is
 125 important that the analyses below are restricted to cells that are truly infected with virus, so we
 126 accepted the possible loss of some low-level infections in order to avoid false positives. In addition,
 127 the synonymous viral barcodes only identify co-infections by viruses with different barcodes – since
 128 the barcodes are at roughly equal proportion, we expect to miss about half of the co-infections.
 129 Since we annotate about ~10% of the infected cells as co-infected by viruses with different barcodes
 130 (Figure 4D), we expect another ~10% of the infected cells to also be co-infected but not annotated
 131 as so by our approach. Because most cells are not infected, we subsampled the uninfected cells to
 132 the numbers shown in Figure 4D to balance the proportions of infected and uninfected cells for all
 133 subsequent analyses.

134 Strikingly, the extreme variation in the number of viral transcripts per cell remains even after we
 135 apply these rigorous criteria for annotating infected cells (Figure 4E). The fraction of viral mRNA per
 136 infected cell follows a roughly exponential distribution, with many cells having few viral transcripts
 137 and a few cells having many. At 6 and 8 hours <10% of infected cells are responsible for over half
 138 the viral transcripts, while at 10 hours <15% of infected cells produce over half the viral transcripts
 139 (Figure 4-Figure supplement 3). One way to quantify the heterogeneity of a distribution is to
 140 calculate the Gini coefficient ([Gini, 1921](#)), which ranges from 0 for a completely uniform distribution,
 141 to 1 for a maximally skewed distribution. Figure 4-Figure supplement 3 shows the Gini coefficients
 142 for the distribution of viral mRNA across infected cells for each sample. The Gini coefficients
 143 are ≥0.64 for all samples. As a fun point of comparison, these Gini coefficients indicate that the
 144 distribution of viral mRNA across infected cells is more uneven than the distribution of income in
 145 the United States ([Alvaredo, 2011](#)).

146 One possible source of heterogeneity in the amount of viral mRNA per cell is variability in the
 147 timing of infection. If some cells are infected earlier in the experiment than others, then they might
 148 have substantially more viral mRNA. However, several lines of evidence indicate that this is not
 149 the major cause of heterogeneity across cells. First, the sample for which the infection inoculum
 150 was never removed (8hr-2) only shows slightly more heterogeneity than samples for which the
 151 inoculum was washed away after one hour (Figure 4E, Figure 4-Figure supplement 3), despite the
 152 fact that the potential time window for infection is much longer in the former sample. Second, in an
 153 independent experiment, we performed completely synchronized infections by pre-binding virus to
 154 cells on ice and then washing away unbound virus before bringing the cells to 37°C [\[Alistair, can you
 155 add relevant citation for this method?\]](#). As shown in Figure 4-Figure supplement 4, flow cytometry
 156 staining found that the heterogeneity in the levels of individual viral proteins was not markedly
 157 different for these synchronized infections than in the absence of pre-binding and washing. Finally,
 158 viral mRNA expression from the secondary spread of virus from infected cells does not appreciably
 159 occur during the timeframes of our experiments, since Figure 4B does not show the pervasive
 160 presence of mixed barcodes that would occur in this case. Therefore, variability in the timing of
 161 infection is not the dominant cause of the cell-to-cell heterogeneity in our experiments.

162 Notably, Figure 4E shows that there are co-infected cells with both low and high amounts of
 163 viral mRNA, suggesting that the initial infectious dose does not drive a simple continuous increase
 164 in viral transcript production. In support of this view, we used flow cytometry to quantify the
 165 levels of individual viral proteins in cells infected at various MOIs or for which we could delineate
 166 co-infection status (Figure 4-Figure supplement 5). This analysis shows that sub-populations of cells
 167 that express similarly low and high levels of viral proteins persist across a wide range of infectious
 168 doses, although co-infection can influence the relative proportion of infected cells that fall into
 169 these sub-populations (Figure 4-Figure supplement 5).

170 **Absence of viral genes partially explains cell-to-cell variability in viral load.**

171 The influenza genome is segmented, and cells can fail to express a viral mRNA if the encoding
 172 gene segment is not packaged in the infecting virion or fails to initiate transcription after infection.
 173 Indeed, several groups have reported that the majority of infected cells fail to express at least
 174 one viral gene (*Brooke et al., 2013; Heldt et al., 2015; Dou et al., 2017*). We wondered if the
 175 absence of specific viral genes might be associated with reduced amounts of viral mRNA within
 176 single infected cells. In particular, transcription of influenza virus mRNAs is performed by the viral
 177 ribonucleoprotein (RNP) complex, which consists of the three proteins that encode the tripartite
 178 polymerase (PB2, PB1, and PA) as well as nucleoprotein (NP) (*Huang et al., 1990*). Each viral gene
 179 segment is associated with one RNP in incoming infecting virions, but secondary transcription by
 180 newly synthesized RNPs requires the presence of the viral genes encoding each of the four RNP
 181 proteins (*Vreede et al., 2004; Eisfeld et al., 2015*). This secondary transcription is a major source
 182 of viral mRNAs, as evidenced by the fact that blocking synthesis of the RNP proteins reduces the
 183 amount of viral mRNA by several orders of magnitude in bulk cells (Figure 5-Figure supplement 1).

184 We examined the total amount of viral mRNA versus the expression of the genes from each
 185 viral segment (Figure 5A, Figure 5-Figure supplement 2, Figure 5-Figure supplement 3). Note
 186 that influenza virus expresses ten major gene transcripts from its eight gene segments, as the
 187 M and NS segments are alternatively spliced to produce the M1 / M2 and NS1 / NEP transcript,
 188 respectively (*Dubois et al., 2014*). However, an inherent limitation of current established single-cell
 189 mRNA sequencing techniques is that they only sequence the 3' end of the transcript (*Zheng et al.,
 190 2017; Macosko et al., 2015; Klein et al., 2015; Cao et al., 2017*). Since the alternative spliceforms
 191 M1 / M2 and NS1 / NEP share the same 3' ends, we cannot distinguish them and therefore will refer
 192 simply to the combined counts of transcripts from each of these alternatively spliced segments as
 193 the M and NS genes.

194 Cells that lack an RNP gene never derive more than a few percent of their mRNAs from virus,
 195 confirming the expected result that all four RNP genes are essential for high levels of viral transcrip-
 196 tion (Figure 5A, Figure 5-Figure supplement 2, Figure 5-Figure supplement 3). However, we observe
 197 cells that lack each of the other non-RNP genes but still derive \approx 40% of their mRNAs from virus,
 198 suggesting that none of the other genes are important for high levels of viral transcription. These re-
 199 sults are statistically supported by Figure 5B, which shows that absence of any RNP gene but *not* any
 200 other viral gene is associated with reduced amounts of viral mRNA. However, gene absence clearly
 201 does not explain all of the variability in viral gene expression, since even cells expressing all viral
 202 genes exhibit a very wide distribution in the amount of viral mRNA that they express. Specifically, at
 203 both 8 and 10 hours, the amount of viral mRNA in individual cells expressing all eight viral genes still
 204 ranges from <1% to >50% (Figure 5A, Figure 5-Figure supplement 2, Figure 5-Figure supplement 3).

205 We also quantified the fraction of infected cells that completely failed to express a given gene. We
 206 limited this analysis to examining the presence / absence of the non-RNP genes in cells expressing
 207 all four RNP genes, since we might fail to detect viral transcripts that are actually present at low
 208 levels in RNP-deficient cells due to the lower viral burden in these cells. At the 8- and 10-hour time
 209 points, between 5% and 17% of cells fail to express any one of the four non-RNP genes (Figure 5C
 210 and Figure 5-source data 1). The absence of a given gene appears to be an independent event, as
 211 the probability of observing all four non-RNP genes in a cell is well predicted by simply multiplying
 212 the probabilities of observing each gene individually (Figure 5C and Figure 5-source data 1). If we
 213 extrapolate the frequencies at which cells lack non-RNP genes to the RNP genes, then we would
 214 predict that 35-50% of infected cells express mRNAs from all eight genes. This estimate of the
 215 frequency at which infected cells express mRNAs from all eight gene segments is slightly higher than
 216 previous estimates of 13% (*Brooke et al., 2013*) and 20% (*Dou et al., 2017*). At least one difference
 217 is that *Brooke et al. (2013)* stained for proteins whereas we examined the expression of mRNAs – it
 218 is likely that some cells contain mutated viral genes that fail to produce stable protein even when
 219 mRNA is expressed.

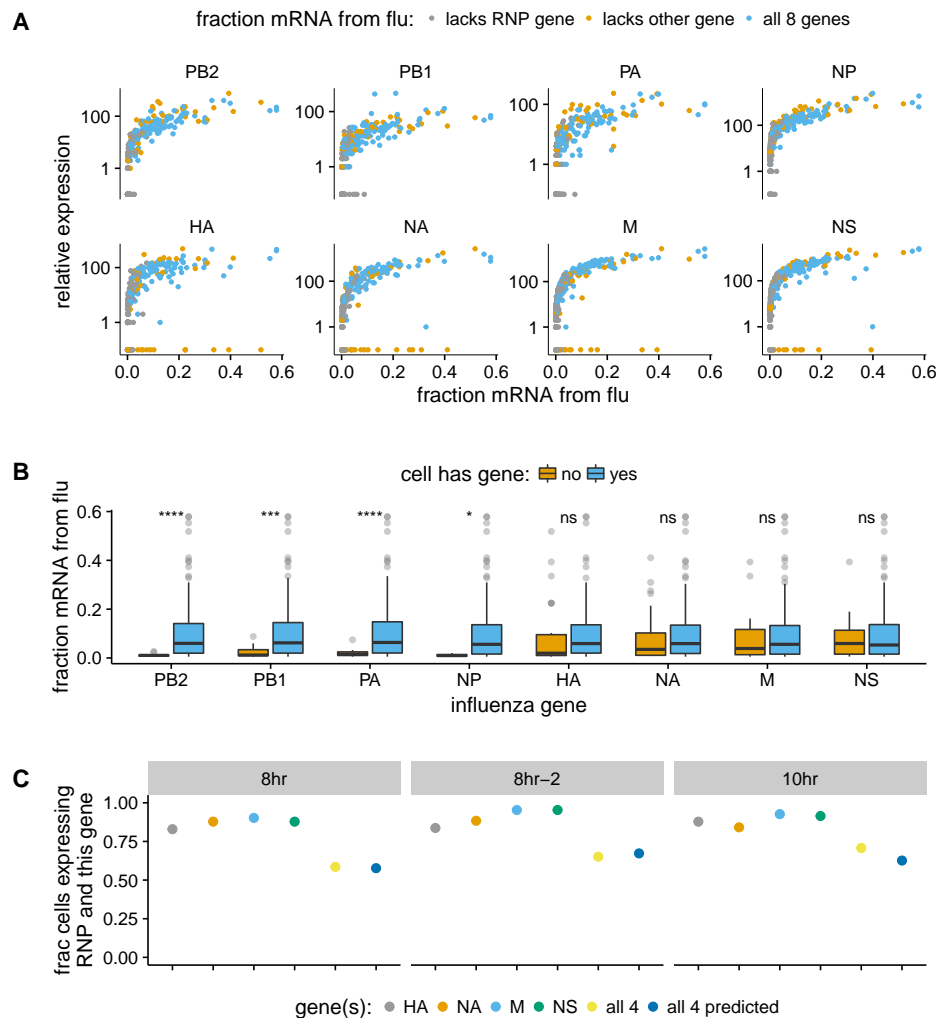


Figure 5. The absence of viral genes explains some of the variability in the amount of viral mRNA per cell. **(A)** The normalized expression of each viral gene as a function of the total fraction of mRNA in each infected cell derived from virus, taken over all time points. Cells with high viral burden always express all RNP genes, but some cells with high viral burden lack each of the other genes. Plots for individual samples are in Figure 5-Figure supplement 2, and a plot that excludes known coinfecting cells is in Figure 5-Figure supplement 3. **(B)** Box and whisker plots showing the per-cell viral burden among cells with >0.5% of their mRNA from virus, binned by whether or not the cells express each gene. A Wilcoxon signed-rank test was used to test the null hypothesis that absence of each gene does not affect viral burden: **** = $P < 10^{-4}$, *** = $P < 10^{-3}$, * = $P < 0.05$, ns = not significant. The trends are similar if we look only at the 10-hour sample (Figure 5-Figure supplement 4) or exclude known co-infected cells (Figure 5-Figure supplement 5). **(C)** The fraction of cells that express each of the four other genes among cells that express all RNP genes, as well as the fraction that express all four of the other genes. The fraction that express all four genes is well predicted by simply multiplying the frequencies of cells that express each gene individually, indicating that gene absence is approximately independent across these genes.

Figure 5-Figure supplement 1. Secondary transcription is a major source of viral mRNA during bulk infections.

Figure 5-Figure supplement 2. Like panel (A), but shows samples individually.

Figure 5-Figure supplement 3. Like panel (A), but excludes coinfecting cells with mixed viral barcodes.

Figure 5-Figure supplement 4. Like panel (B) but for the 10-hr sample only.

Figure 5-Figure supplement 5. Like panel (B) but excludes coinfecting cells with mixed viral barcodes.

Figure 5-source data 1. The numerical data for panel (C) are in p_missing_genes.csv.

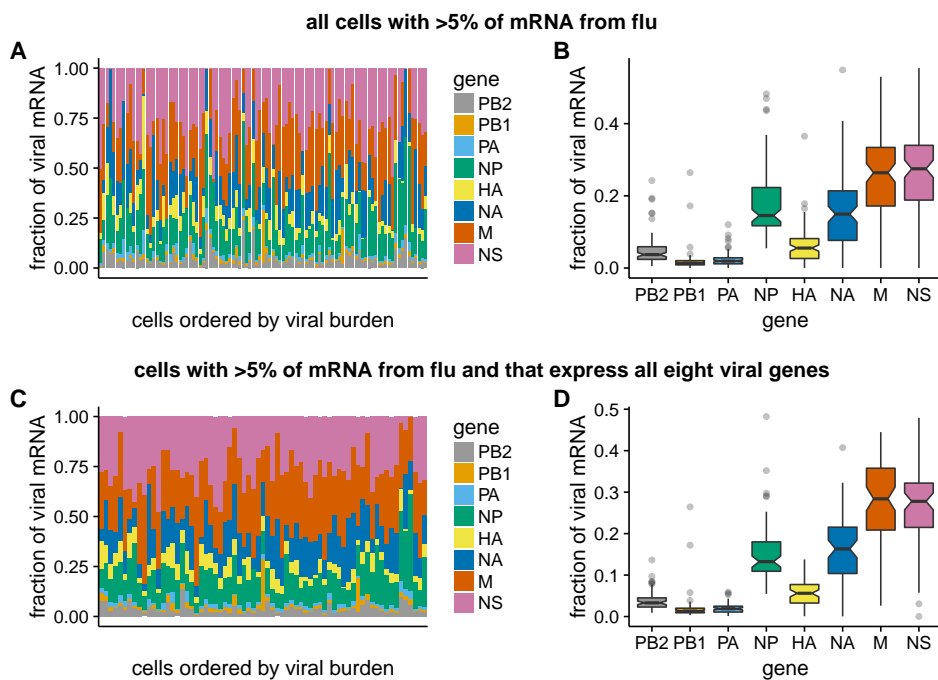


Figure 6. [add expression wedges] Relative expression of influenza virus genes in highly infected cells (>5% of total mRNA from virus). **(A)** The fraction of viral mRNA from each viral gene for each cell. **(B)** Box plots showing the distribution of the fraction of viral mRNA per cell from each viral gene. The black lines at the notches are the medians, and the tops and bottoms of boxes indicate the first and third quartiles. Whiskers extend to the highest or lowest data point observed within 1.5x the interquartile range, outliers shown as circles. Notches extend 1.58x the interquartile range divided by the square root of the number of observations. **(C), (D)** The same plots, but only including cells for which we observed at least one molecule of each viral gene.

Figure 6-source data 1. The raw data for all cells are in `p_flu_expr_all.csv`.

Figure 6-source data 2. The raw data for fully infected cells are in `p_flu_expr_fullyinfected.csv`.

220 **The relative amounts of different viral mRNAs are more consistent across cells.**
 221 The results above show that the amount of viral mRNA in infected cells varies over several orders
 222 of magnitude. Does the relative expression of viral genes exhibit similar cell-to-cell variability?
 223 To address this question, we focused on cells that derived >5% of their mRNA from virus, since
 224 estimates of relative viral gene expression will be less noisy in cells with more viral mRNAs.

225 In contrast to the extreme variability in the total viral mRNA per cell, the fraction of this mRNA
 226 derived from each gene is much more consistent across cells (Figure 6A). Total viral mRNA varies by
 227 orders of magnitude, but the fraction from any given viral gene is fairly tightly clustered around the
 228 median value for all cells (Figure 6B). The relative levels of each viral mRNA in our cells are similar
 229 to prior bulk measurements made by Northern blots (*Hatada et al., 1989*), which also found an
 230 expression hierarchy of M > NS > NP > NA > HA > PB2 ~ PB1 ~ PA. The cell-to-cell consistency in
 231 the relative expression of different viral genes is even tighter if we limit the analysis only to cells
 232 that express all eight viral genes (Figure 6C,D). Therefore, with the exception of complete gene
 233 absence, the factors that drive the dramatic cell-to-cell variability in the amount of viral mRNA have
 234 roughly similar effects on all viral genes in a given cell. This finding is consistent with prior work
 235 showing positive correlations among the abundance of several viral genome segments in individual
 236 cells (*Heldt et al., 2015*).

237 **Co-infection can provide infected cells with the full complement of viral genes.**

238 Our sequencing enables us to identify the rare cells that were co-infected with both wild-type and
 239 synonymous barcoded viral variants. Overall, we captured 10 such co-infected cells that had >5%

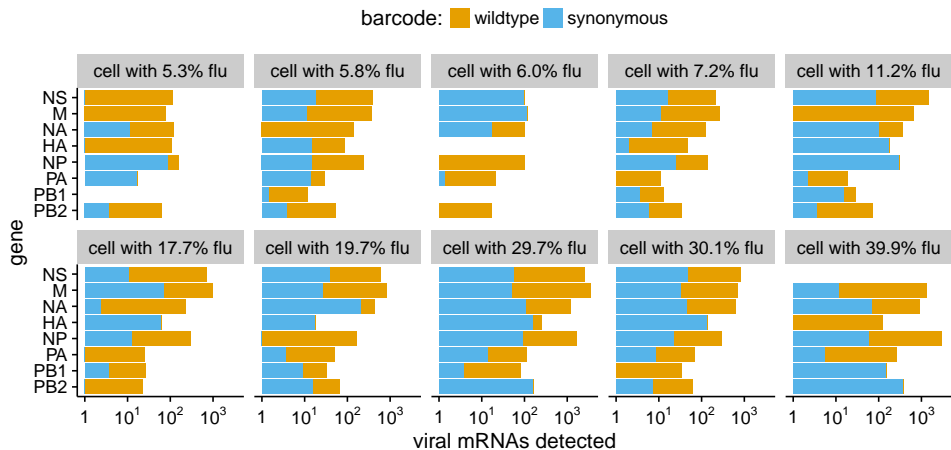


Figure 7. The abundance of each viral transcript in cells that are co-infected with the two viral variants and have >5% of their mRNA derived from virus. The bars show the logarithms of the numbers of each viral mRNA detected, and are colored in linear proportion to the fraction of that mRNAs derived from wild-type or synonymously barcoded virus.

Figure 7-Figure supplement 1. Co-infected cells express roughly equal amounts of a gene from each infecting viral variant.

Figure 7-source data 1. The raw data plotted in this figure are in `p_co-infection.csv`.

Figure 7-source data 2. The sequence of the HA viral RNA carrying the GFP gene is in `HAflank-eGFP.fasta`.

of their mRNA derived from virus (Figure 7). Seven of these 10 cells expressed all eight viral genes. The majority (4 of 7) of these cells would *not* have expressed all the viral genes in the absence of co-infection, since they have at least one gene exclusively derived from each viral variant. For instance, the cell with 11.2% of its mRNA from virus in the upper right of Figure 7 expresses M only from the wildtype viral variant, and NP and HA only from the synonymously barcoded variant. Our data therefore provide the first direct single-cell observation of the fact that co-infection can rescue missing viral genes (*Brooke et al., 2013, 2014; Fonville et al., 2015; Aguilera et al., 2017*).

Another observation from Figure 7 is that co-infected cells usually express roughly equal amounts of transcripts from each of the two viral variants. This observation is consistent with the finding by *Dou et al. (2017)* and *Huang et al. (2008)* that the temporal window for co-infection is short – if both viral variants infect a cell at about the same time, then neither will have a headstart and so each will have a roughly equal opportunity to transcribe its genes.

To support this idea with a larger dataset albeit at lower resolution, we generated a virus in which the HA coding sequence was replaced by GFP. We then co-infected cells with a mix of wildtype and ΔHA-GFP virus and used flow cytometry to score cells for the presence of HA only (infection by wildtype virus), GFP only (infection by ΔHA-GFP virus), or both (co-infection) as shown in Figure 7 Figure supplement 1. As in our single-cell sequencing data, we found that expression of HA and GFP were highly correlated, indicating that co-infected cells typically expressed roughly equal amounts of transcript from each viral variant.

Activation of the interferon response is rare in single infected cells.

Because our sequencing captured all polyadenylated transcripts, we can examine whether there are prominent changes in the host-cell transcriptome in sub-populations of infected cells. Influenza virus infection can trigger innate-immune sensors that lead to the transcriptional induction of type I and III interferons, and subsequently of anti-viral interferon-stimulated genes (*Killip et al., 2015; Iwasaki and Pillai, 2014; Crotta et al., 2013*). However, activation of the interferon response is stochastic and bi-modal at the level of single cells (*Chen et al., 2010; Shalek et al., 2013, 2014; Perez-Cidoncha et al., 2014; Bhushal et al., 2017; Hagai et al., 2017*). We therefore hypothesized

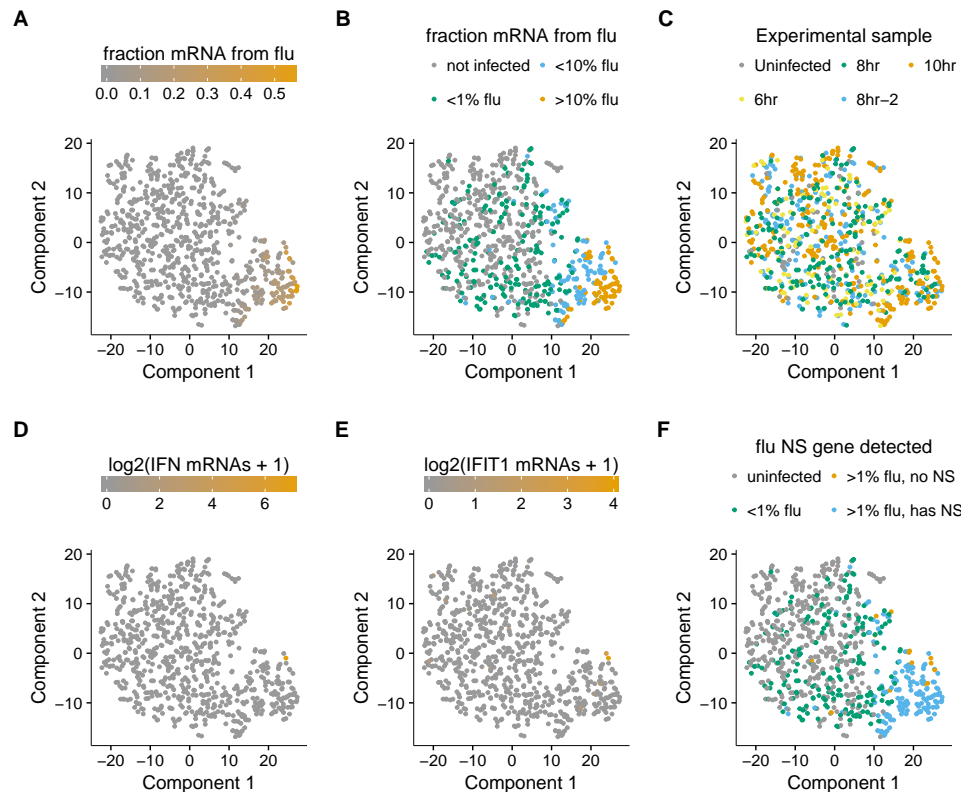


Figure 8. [add arrow] A t-SNE plot created by semi-supervised clustering using genes that co-vary with viral infection status. Each point is a single cell, and each panel shows an identical layout but colors the cells according to a different property. **(A), (B)** Cells colored by the fraction of their mRNA derived from virus. **(C)** Cells colored by the experimental sample. **(D), (E)** Cells colored by the number of detected transcripts from type I and III interferons (IFN). Only one cell has detectable interferon expression (in orange, indicated with arrow). **(E)** Cells colored by the expression of the interferon-stimulated gene IFIT1. **(F)** Cells colored by whether they express the viral NS gene. The one interferon-positive cell is lacking NS, but so are many interferon-negative cells.

267 that we might see two sub-populations of infected cells: one in which the interferon response
268 inhibited viral transcription, and another in which the virus was able to express high levels of its
269 mRNA by evading or blocking this response.

270 To examine whether there were distinct sub-populations of virus-infected cells, we used a
271 semi-supervised t-SNE approach (*Van der Maaten and Hinton, 2008*) to cluster cells by genes that
272 co-varied with viral infection status. As shown in Figure 8A,B, this approach effectively grouped cells
273 by the amount of viral mRNA that they expressed. Sample-to-sample variation was regressed away
274 during the clustering, as cells did not obviously group by time-point, with expected exception that
275 the uninfected and 6-hour samples had few cells in the region of the plot corresponding to large
276 amounts of viral mRNA (Figure 8C).

277 But to our surprise, we did not see a prominent clustering of infected cells into sub-populations
278 as expected if the interferon response was strongly activated in some cells. To investigate fur-
279 ther, we annotated each cell by the total number of type I and III interferon transcripts detected.
280 Remarkably, only a single cell expressed detectable interferon (Figure 8D). We also examined
281 interferon-stimulated genes, which are induced by autocrine and paracrine interferon signaling.
282 Figure 8E shows expression of one such gene, IFIT1 (*Fensterl and Sen, 2011*). As with interferon
283 itself, expression of IFIT1 was rare and most prominent in the single interferon-positive cell, pre-
284 sumably due to the higher efficiency of autocrine versus paracrine signaling. Notably, interferon

285 and interferon-stimulated genes were also relatively ineffective at blocking viral transcription in the
 286 single cell in which they were potently induced, since >10% of the mRNA in this cell was derived
 287 from virus (Figure 8A,B,D,E).

288 We posited that the paucity of interferon induction might be due to the activity of influenza
 289 virus's major interferon antagonist, the NS1 protein (*García-Sastre et al., 1998; Hale et al., 2008*).
 290 We therefore identified cells that expressed substantial amounts of viral mRNA but lacked the
 291 NS gene (Figure 8F). Consistent with the idea that NS1 is important for suppressing interferon,
 292 the one interferon-positive cell lacked detectable expression of the NS gene. But other cells that
 293 lacked NS expression still failed to induce a detectable interferon response, despite often having a
 294 substantial amount of their mRNA derived from virus (Figure 8). This result is in line with other work
 295 showing that NS1-deficient influenza virus does not deterministically induce interferon (*Killip et al.,
 296 2017; Kallfass et al., 2013*). Therefore, many individual infected cells fail to activate innate-immune
 297 responses even when the virus lacks its major interferon antagonist.

298 **Some host genes co-vary with viral gene expression.**

299 We examined whether any host genes were differentially expressed in cells with more viral mRNA.
 300 We restricted this analysis to infected cells with all eight viral genes in order to focus on cellular
 301 genes that were associated with viral mRNA burden independent of effects due to the presence
 302 or absence of particular viral transcripts. We identified 43 cellular genes that co-varied with viral
 303 mRNA expression at a false discovery rate of 0.1 (Figure 9, Figure 9-source data 1).

304 A gene-set analysis shows that many cellular genes that are associated with the amount of viral
 305 mRNA are involved in the response to reactive oxygen species or hypoxia (Figure 9-Figure supple-
 306 ment 2). Genes known or suspected to be regulated by the Nrf2 master regulator in response to
 307 oxidative stress are often expressed at higher levels in cells with more viral mRNA (Figure 9). These
 308 genes produce proteins that are involved in detoxification of reactive oxygen species or resultant
 309 products, the management of misfolded proteins, the electron transport chain, or a general stress
 310 response (Figure 9-Figure supplement 1). We additionally see reduced expression of the nitric
 311 oxide synthase interacting protein (NOSIP). Transient oxidative stress is known to occur during viral
 312 infection, and may act in a proviral fashion via MAPK activation driving vRNP export (*Amatore et al.,
 313 2014*). The antioxidant response is thought to be largely antiviral, potentially through inhibition of
 314 MAPK activity (*Lin et al., 2016; Sgarbanti et al., 2014*). To directly test the effect of transient oxida-
 315 tive stress, we compared the fraction of cells that expressed detectable viral protein when infected
 316 either with or without pre-treatment to induce oxidative stress. Figure 9-Figure supplement 2 shows
 317 that the cells pre-treated to induce oxidative stress exhibited more frequent detectable expression
 318 of viral protein. These results, in conjunction with the differential expression test in Figure 9 and
 319 the prior work mentioned above, suggest that oxidative stress acts in a proviral fashion.

320 The gene-set analysis also found that the amount of viral mRNA was associated with the
 321 expression of genes involved in the G2-M cell-cycle checkpoint (Figure 9-Figure supplement 2). The
 322 cell-cycle associated genes CCND3, MKI67, UBE2S, and CENPF are all expressed at significantly lower
 323 levels in cells with more viral mRNA (Figure 9). However, our data are not sufficient to determine
 324 whether the lower expression of these genes is a cause or effect of the reduction in viral mRNA.

325 Interestingly, none of the cellular genes that are significantly associated with the amount of viral
 326 mRNA in our study are among the 128 genes that *Watanabe et al. (2010)* report as having been
 327 identified multiple times in genome-wide screens for factors affecting influenza virus replication.
 328 One possible explanation is that most of the cell-to-cell heterogeneity in our experiments might
 329 arise from viral factors, pure stochasticity, or more subtle alterations in host-cell state – not due
 330 to changes of expression in the type of single large-effect genes that are usually identified in
 331 genome-wide knockdown / knockout studies.

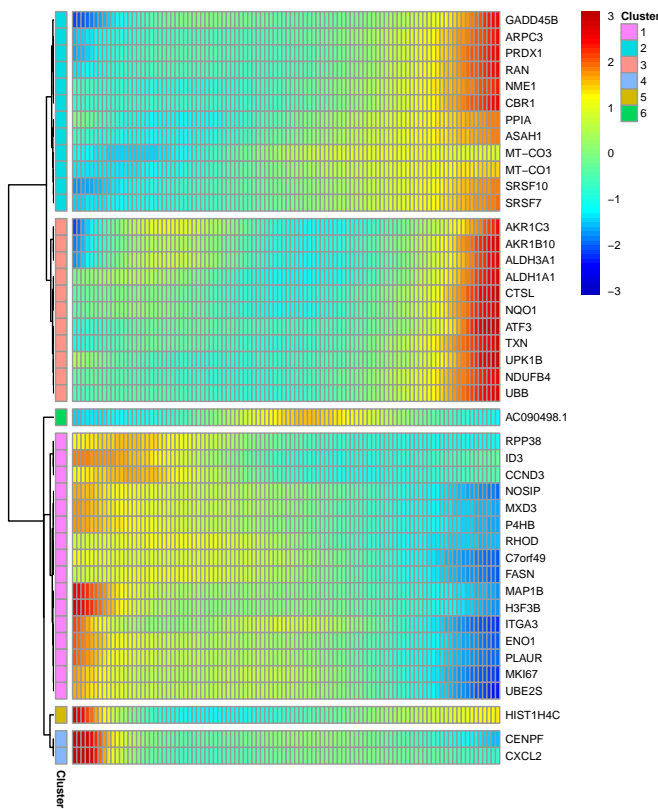


Figure 9. [add gray arrow wedge] Cellular genes that co-vary in expression with the amount of viral mRNA in cells expressing all eight viral genes. The columns are cells, ordered from left to right by the fraction of mRNA derived from virus. Each row is a gene that is differentially expressed as a function of the fraction of mRNA derived from virus at a false discovery rate of 0.1. Genes for which the color goes from blue at left to red at right are expressed at higher levels in cells with more viral mRNA. The scale bar indicates the number of standard deviations above or below the mean expression, truncated at 3-fold on both sides.

Figure 9-source data 1. The full results of the differential expression test are in p_sig_cellular_genes.csv.

Figure 9-source data 2. A gene-set analysis for pathways associated with the amount of viral mRNA is in p_pathway_enrichment.csv.

Figure 9-Figure supplement 1. Many genes that co-vary with viral load are involved in the oxidative stress response.

Figure 9-Figure supplement 2. Pre-treating to induce oxidative stress increases the fraction of infected cells expressing detectable viral protein.

Discussion

We have quantified the total transcriptome composition of single cells infected with influenza virus. While we observe a general increase in the amount of viral mRNA over time as expected from bulk measurements (*Hatada et al., 1989; Shapiro et al., 1987*), there is wide variation in viral gene expression among individual infected cells.

The most obvious form of heterogeneity is the complete failure of some infected cells to express one or more viral genes, which we estimate occurs in about half the infected cells in our experiments. The absence of some viral genes in some infected cells has been noted previously (*Brooke et al., 2013; Heldt et al., 2015; Dou et al., 2017*), and our work provides a holistic view by quantifying the total viral transcriptional load as a function of the level of each mRNA. We find that cells lacking expression of any of the four genes that encode the viral RNP express much less total viral mRNA, consistent with prior bulk studies (*Vreede et al., 2004; Eisfeld et al., 2015*). Interestingly, the reason some cells fail to express some viral genes remains unclear. The prototypical influenza virion packages one copy of each of the eight gene segments (*Noda et al., 2006; Hutchinson et al., 2010*),

346 but some virions surely package fewer (*Brooke et al., 2014*). However, it is also possible that much
 347 of the viral gene absence is due to stochastic loss of viral RNPs after infection but prior to the
 348 initiation of viral transcription in the nucleus.

349 The absence of viral genes only partially explains the cell-to-cell variation in amount of viral
 350 mRNA, which still varies from <1% to >50% among cells expressing all the viral genes. It is likely
 351 that other viral genetic factors explain some of this remaining heterogeneity. The 3'-end sequencing
 352 strategy used in our experiments detects the presence of a viral gene, but does not identify
 353 whether that gene contains a mutation that might hinder viral replication. However, viral mutations
 354 are also unlikely to explain all the observed heterogeneity, since current consensus estimates of
 355 influenza virus's mutation rate suggest that the typical virion in a stock such as the one used in our
 356 experiment should contain less than one mutation per genome (*Parvin et al., 1986; Suárez et al.,
 357 1992; Suárez-López and Ortín, 1994; Nobusawa and Sato, 2006; Bloom, 2014; Pauly et al., 2017*).

358 The rest of the heterogeneity must be due to some combination of cellular factors and inherent
 359 stochasticity. Some features of the cellular transcriptome co-vary with the amount of influenza
 360 mRNA. In particular, the viral load in individual cells is associated with the expression of genes
 361 involved in response to cellular stresses, including oxidative stress. It will be interesting to determine
 362 if these cellular transcriptional signatures are simply a consequence of the stress imposed by viral
 363 replication, or if their stronger activation in some cells is a causative factor that promotes viral
 364 transcription. However, it also would not be surprising if a substantial amount of the cell-to-cell
 365 heterogeneity cannot be ascribed to pre-existing features of either the viral genome or cellular state.
 366 Apparently stochastic heterogeneity is a common feature of many processes at a single-cell level (*Cai
 367 et al., 2006; Raj et al., 2006; Buganim et al., 2012; Shalek et al., 2013; Avraham et al., 2015*) –
 368 especially when those processes are initiated by very small numbers of initial molecules (*Elowitz
 369 et al., 2002*), as is the case for low-MOI viral infection.

370 Our data do suggest that the factors driving the heterogeneity in viral transcriptional load exert
 371 relatively concordant effects on all viral genes in a given cell. Specifically, despite the extreme
 372 heterogeneity in total viral mRNA per cell, the relative levels of the viral mRNAs are reasonably
 373 consistent across cells, and generally reflective of classical bulk measurements (*Hatada et al., 1989*).
 374 Therefore, despite the stochasticity inherent in initiating transcription and replication of each gene
 375 from a single copy carried by the incoming virion, as long as a gene is not completely lost then the
 376 virus possesses mechanisms to control its relative expression (*Shapiro et al., 1987; Hatada et al.,
 377 1989; Perez et al., 2010; Heldt et al., 2012; Chua et al., 2013*).

378 One factor that surprisingly does *not* appreciably contribute to the heterogeneity in our ex-
 379 periments is activation of innate-immune interferon pathways. Only one of the hundreds of
 380 virus-infected cells expresses any detectable interferon, despite the fact that a number of cells fail
 381 to express the influenza-virus interferon antagonist NS1. It is known that interferon activation is
 382 stochastic at the level of single cells in response to both synthetic ligands (*Shalek et al., 2013, 2014;
 383 Bhushal et al., 2017; Hagai et al., 2017*) and actual infection (*Rand et al., 2012; Perez-Cidoncha
 384 et al., 2014; Avraham et al., 2015; Killip et al., 2017*). But interferon expression is a prominent
 385 transcriptional signature of high-MOI influenza virus infection of bulk cells, including in the epithelial
 386 cell line and at the time-points used in our experiments (*Geiss et al., 2002; Sutejo et al., 2012*). So
 387 it is notable how rarely single cells express interferon. Interferon expression would surely be more
 388 common at later times or with a viral stock passaged at higher MOI, since paracrine interferon
 389 signaling (*Crotta et al., 2013*) and accumulation of defective viral particles enhance innate-immune
 390 detection (*Tapia et al., 2013; Lopez, 2014*). However, the early events of physiological influenza
 391 infection involve just a few virions (*Varble et al., 2014; McCrone et al., 2017*), and so it is interesting
 392 to speculate whether rare events such as interferon activation during the first few cycles of viral
 393 replication could contribute to heterogeneity in the eventual outcome of infection.

394 Overall, our work shows the power and importance of characterizing cellular infection at the
 395 level of single cells (*Avraham et al., 2015*). The dynamics of viral infection in any given cell is shaped
 396 by the genetic composition of the incoming virion, the host-cell state, the bi-modality of innate-

397 immune activation, and the inherent stochasticity of molecular processes initiated by a single copy
 398 of each viral gene. We have shown how the confluence of these factors leads to extreme cell-to-cell
 399 heterogeneity in the transcriptional outcome of influenza virus infection. Further deconstruction of
 400 the contributions of each factor will enable a deeper understanding of how the bulk features of
 401 infection emerge from the processes occurring within individual virus-infected cells.

402 Methods and Materials

403 Cell lines and viruses

404 The following cell lines were used in this study: the human lung epithelial carcinoma line A549
 405 (ATCC CCL-185), the MDCK-SIAT1 variant of the Madin Darby canine kidney cell line overexpressing
 406 human SIAT1 (Sigma-Aldrich 05071502), and the human embryonic kidney cell line 293T (ATCC
 407 CRL-3216). All cells were maintained in D10 media (DMEM supplemented with 10% heat-inactivated
 408 fetal bovine serum, 2 mM L-glutamine, 100 U of penicillin/ml, and 100 µg of streptomycin/ml) at 37
 409 °C at 5 % CO₂.

410 Wildtype A/WSN/1933 (H1N1) influenza virus was generated by reverse genetics using the
 411 plasmids pHW181-PB2, pHW182-PB1, pHW183-PA, pHW184-HA, pHW185-NP, pHW186-NA, pHW187-
 412 M, and pHW188-NS (**Hoffmann et al., 2000**). The sequences of the viral RNAs encoded in these
 413 plasmids are in Figure 1-source data 1. Reverse-genetics plasmids encoding the synonymously
 414 barcoded WSN virus were created by using site-directed mutagenesis to introduce two synonymous
 415 mutations near the 3' end of the mRNA for each viral gene. The sequences of the synonymously
 416 barcoded viral RNAs are in Figure 1-source data 1.

417 To generate viruses from these plasmids, we transfected an equimolar mix of all eight plasmids
 418 into cocultures of 293T and MDCK-SIAT1 cells seeded at a ratio of 8:1. At 24 hours post-transfection,
 419 we changed media from D10 to influenza growth media (Opti-MEM supplemented with 0.01% heat-
 420 inactivated FBS, 0.3% BSA, 100 U of penicillin/ml, 100 µg of streptomycin/ml, and 100 µg of calcium
 421 chloride/ml). At 48 hours post-transfection we harvested the virus-containing supernatant, pelleted
 422 cellular material by centrifugation at 300 x g's for 4 minutes, and stored aliquots of the clarified
 423 viral supernatant at -80 °C. We then titered thawed aliquots of viral by TCID50 on MDCK-SIAT1 cells,
 424 computing titers via the formula of **Reed and Muench (1938)**. To generate our "high-purity" stocks
 425 of viruses for the single-cell sequencing experiments, we then infected MDCK-SIAT1 cells at an MOI
 426 of 0.01, and let the virus grow for 36 hours prior to harvesting aliquots that were again clarified by
 427 low-speed centrifugation, aliquoted, stored at -80 °C, and titered by TCID50. The high-MOI passage
 428 (high-defective particle) stock used in Figure 2 was generated by instead passaging in MDCK-SIAT1
 429 cells twice at an MOI of 1 for 48 hours.

430 For the experiments in Figure 7-Figure supplement 1, we created a virus that carried an HA gene
 431 segment in which GFP replaced most of the HA coding sequence, following a scheme first described
 432 by **Marsh et al. (2007)**. Briefly, we created a plasmid encoding a viral RNA with GFP in place of the
 433 HA coding sequence in the context of the pH21 (**Neumann et al., 1999**) reverse-genetics plasmid,
 434 removing potential start codons upstream of the GFP (see Figure 7-source data 2 for the sequence
 435 of the viral RNA). We then generated GFP-carrying virus by reverse-genetics in cells constitutively
 436 expressing HA (**Doud and Bloom, 2016**). To obtain sufficient titers, this HA-eGFP virus was expanded
 437 for 44 rather than 36 hours after initiating infection at an MOI of 0.01.

438 qPCR

439 For the qPCR in Figure 2 and Figure 5-Figure supplement 1, A549 cells were seeded at 3x10⁵
 440 cells per well in a 6-well tissue culture plate in D10 the day prior to infection. On the day of
 441 infection, a single well was trypsinized and the cells were counted in order to determine the
 442 appropriate amount of virus to use to achieve the intended MOI. Immediately before infection,
 443 D10 was replaced with influenza growth media. For cells incubated with cyclohexamide, the
 444 compound was added to a final concentration of 50 µg/ml at the time of infection – previously

445 confirmed to be sufficient to halt viral protein production (*Killip et al., 2014*). RNA was purified
 446 using the QIAGEN RNeasy plus mini kit following manufacturer's instructions. cDNA was syn-
 447 thesized using an oligoDT primer and the SuperScript™ III first-strand synthesis supermix from
 448 ThermoFisher using the manufacturer's protocol. Transcript abundance was measured using
 449 SYBR™ green PCR master mix, using a combined anneal/extension step of 60 °C for one minute
 450 with the following primers: *HA*: 5'-GGCCCAACCACACATTCAAC-3', 5'-GCTCATCACTGCTAGACGGG-
 451 3', *IFNB1*: 5'-AAACTCATGAGCAGTCTGCA-3', 5'-AGGAGATCTTCAGTTCCGGAGG-3', *L32*: 5'-
 452 AGCTCCAAAAATAGACGCAC-3', 5'-TTCATAGCAGTAGGCACAAAGG-3'. Biological triplicates were per-
 453 formed for all samples.

454 For the measurements of viral genomic HA content in Figure 2A, vRNA was harvested from 80
 455 µl of viral supernatant by the addition of 600 µl of RLT plus before proceeding with the standard
 456 QIAGEN RNeasy Plus Mini kit protocol. The cDNA was generated using SuperScript™ III first-strand
 457 synthesis supermix using the manufacturer's protocol, and using the universal vRNA primers of
 458 *Hoffmann et al. (2001)* with the modifications described in *Xue et al. (2017)*. The qPCR was then
 459 performed as for mRNA measurements. A standard curve was generated from three independent
 460 dilutions of the HA-encoding reverse genetics plasmid. All vRNA values represent three independent
 461 RNA extractions with two replicate qPCR measurements.

462 **Flow cytometry titering and analyses**

463 To determine viral titers in terms of HA-expressing units and for the flow cytometry in Figure 7,
 464 Figure supplement 1, and Figure supplement 5 A549 cells were seeded in a 6-well plate and infected
 465 as described above for the qPCR analyses. Cells were harvested by trypsinization, resuspended in
 466 phosphate-buffered saline supplemented with 2% heat-inactivated FBS, and stained with 10 µg/ml
 467 of H17-L19, a mouse monoclonal antibody confirmed to bind to WSN HA in a prior study (*Doud*
 468 *et al., 2017*). After washing in PBS supplemented with 2% FBS, the cells were stained with a goat
 469 anti-mouse IgG antibody conjugated to APC. Cells were then washed, fixed in 1% formaldehyde,
 470 and washed further before a final resuspension and analysis. We then determined the fraction of
 471 cells that were HA positive and calculated the HA-expressing units. For NS1 staining in Figure sup-
 472 plement 5, cells stained for HA as described above were permeabilized using BD Cytofix/Cytoperm
 473 following manufacturer's instructions, stained with anti-NS1 (GTX125990, Genetex) at 4.4 µg/ml,
 474 washed, stained with a goat anti-rabbit IgG antibody conjugated to Alexa Fluor 405. washed, and
 475 analyzed. For Figure 7, Figure supplement 1, and Figure supplement 5), after gating to exclude
 476 multiplets in FlowJo, data were extracted using the R package flowCore (*Le Meur et al., 2007*) and
 477 analyzed using a custom Python script. For Figure supplement 5) channels were additionally com-
 478 pensated in FlowJo. Guassian kernel density estimates were obtained using the scipy stats package
 479 method, guassian_kde, using automatic bandwidth determination (*van der Walt et al., 2017*). For
 480 Figure supplement 5), the percentage of influenza-infected cells was determined by HA staining
 481 alone, and the top quantile of NS1-stained cells matching that percentage were taken as the NS1
 482 positive population.

483 **Infections for single-cell mRNA sequencing**

484 Single-cell sequencing libraries were generated using the 10x Chromium Single Cell 3' plat-
 485 form (*Zheng et al., 2017*) using the V1 reagents.

486 All time points except for the second 8-hour sample (8hr-2) were prepared on the same day.
 487 For the infections, A549 cells were seeded in a 6-well plate, with two wells per time point. A single
 488 well of cells was trypsinized and counted prior to initiation of the experiment for the purposes of
 489 calculating MOI. Wild-type and synonymously barcoded virus were mixed to an estimated ratio
 490 of 1:1 based on prior, exploratory, single-cell analyses (data not shown). At the initiation of our
 491 experiment, the wells for all time points were changed from D10 to influenza growth media. Cells
 492 were then infected with 0.3 HA-expressing units of virus per cell (a determined by flow cytometry).
 493 The infections were performed in order of time point: first the 10-hour time point, then the

494 8-hour, and then the 6-hour time point. At one hour after infection, the media for each time
 495 point was changed to fresh influenza growth media. Note that the HA-expressing units were
 496 calculated without this additional washing step, and so likely represent an overestimate of our
 497 final infectious dose (consistent with the fact that fewer than 30% of cells appear infected in the
 498 single-cell sequencing data). All cells were then harvested for single-cell analysis concurrently –
 499 ensuring all had spent equivalent time in changed media . For 8hr-2 sample, cells were infected
 500 as above except that the cells were infected at 0.1 HA-expressing units of virus per cell but no
 501 wash step was performed, and the sample was prepared on a different day. After harvest, cells
 502 were counted using disposable hemocytometers and diluted to equivalent concentrations with an
 503 intended capture of 3000 cells/sample following the manufacturer's provided by 10x Genomics for
 504 the Chromium Single Cell platform. All subsequent steps through library preparation followed the
 505 manufacturer's protocol. Samples were sequenced on an Illumina HiSeq.

506 **Computational analysis of single-cell mRNA sequencing data**

507 Jupyter notebooks that perform all of the computational analyses are available in Supplemen-
 508 tary file 1 and at https://github.com/jbloomlab/flu_single_cell [*This GitHub repository will be made
 509 public upon acceptance of the manuscript. If you are a reviewer who needs access, please contact the
 510 editor.*].

511 Briefly, the raw deep sequencing data were processed using the 10X Genomics software pack-
 512 age CellRanger (version 2.0.0). The reads were aligned to a concatenation of the human and
 513 influenza virus transcriptomes. The human transcriptome was generated by filtering genome
 514 assembly GRCh38 for protein coding genes defined in the GTF file GRCh38.87. The influenza
 515 virus transcriptome was generated from the reverse-complement of the wildtype WSN viral
 516 RNA sequences as encoded in the reverse-genetics plasmids (Figure 1-source data 1). The
 517 aligned deep sequencing data are available on the GEO repository under accession GSE108041
 518 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE108041>).

519 CellRanger calls cells based on the number of observed cell barcodes, and creates a cell-gene
 520 matrix. We used custom Python code to annotate the cells in this matrix by the number of viral
 521 reads that could be assigned to the wildtype and synonymously barcoded virus. Only about half of
 522 the viral reads overlapped the barcoded regions of the genes (Figure 1A) and could therefore be
 523 assigned to a viral barcode (Figure 4-Figures supplement 1). So for calculations of the number of
 524 reads in a cell derived from each viral barcode for each viral gene, the total number of detected
 525 molecules of that gene are multiplied by the fraction of those molecules with assignable barcodes
 526 that are assigned to that barcode. This annotated cell-gene matrix is in Supplementary file 2. A
 527 Jupyter notebook that performs these analyses is in Supplementary file 1.

528 The annotated cell-gene matrix was analyzed in R, primarily using the Monocle package (version
 529 2.4.0) (Qiu et al., 2017; Trapnell et al., 2014). A Jupyter notebook that performs these analyses is in
 530 Supplementary file 1. For each sample, cell barcodes that had >2.5-fold fewer or more UMI counts
 531 mapping to cellular transcripts than the sample mean were excluded from downstream analyses
 532 (see red vertical lines in Figure 3B).

533 In order to determine an appropriate cutoff for how many reads a cell needed to contain in
 534 order to be classified as infected, we calculated the mean viral barcode purity across all cells that
 535 contained at least a given fraction of viral mRNA and had multiple viral reads that could be assigned
 536 a barcode (Figure 4B,C and Figure 4-Figure supplement 2). We then determined the threshold
 537 fraction of viral mRNA at which the mean purity no longer increased as a function of the amount
 538 of viral mRNA. This threshold represents the point at which we have effectively eliminated cells
 539 that have low barcode purity simply due to lysis-acquired reads sampled randomly from both viral
 540 barcodes. As is apparent from Figure 4B, only the 10-hour sample and the 8hr-2 sample have the
 541 excess of mixed barcodes among cells with low amounts of viral mRNA. The likely reason is that
 542 these samples have more total viral mRNA (and so there is more available mRNA to be acquired
 543 from lysed cells); in addition, there is always some experimental variability in the amount of cell

lysis during the 10X sequencing process, and these samples may simply have the most. So the above threshold procedure is appropriate for those two samples. For the other samples, we simply set a minimum threshold of requiring at least a fraction 2×10^{-4} reads to come from viral mRNA as explained in the legend to Figure 4-Figure supplement 2. The thresholds for each sample are shown in Figure 4C and Figure 4-Figure supplement 2. This procedure is expected to be conservative, and may miss some truly infected cells with very low amounts of viral mRNA. For subsequent analyses, we retained all infected cells and a subsample of uninfected cells (the greater of 50 or the number of infected cells for that sample). The rationale for subsampling the uninfected cell is that the vast majority of cells are uninfected, and we did not want these cells to completely dominate the downstream analyses. Cells were classified as co-infected if both viral variants had an RNA level that exceeded the threshold, and if the minor variant contributed at least 5% of the viral mRNA.

For the semi-supervised t-SNE clustering, we used Monocle's cell hierarchy function to bin cells into those with no viral mRNA, <2% viral mRNA, between 2% and 20% viral mRNA, and >20%. Candidate marker genes for t-SNE dimensionality reduction were then determined using the Monocle function markerDiffTable, excluding the effects of sample variation and the number of genes identified in a given cell, using a q-value cutoff of 0.01. The specificity of these markers was determined using the function calculateMarkerSpecificity – the top 50 markers were retained, and used to place populations in a two-dimensional plane based on tSNE dimensionality reduction.

For the analyses of cellular genes that differed in expression as a function of the amount of viral mRNA, we only considered cells that expressed all 8 viral mRNAs to avoid effects driven simply by viral gene absence. We also only considered cellular genes in the differential gene analysis, since viral gene expression will tautologically co-vary with the amount of viral mRNA. Additionally, because influenza virus has the capacity to degrade or prevent the synthesis of host mRNAs (*Bercovich-Kinori et al., 2016*) and contributes significantly to the total number UMIs in some cells, we calculate size factors (a scalar value representing efficiency of UMI capture) based on cellular transcripts alone. Finally, we assigned all cells a ceiling fraction of mRNA from virus of 25% so that a few extremely high-expressing cells did not dominate. Cellular genes with expression that co-varied with the fraction of viral mRNAs in a cell were then determined using the Monocle differentialGeneTest, after removing variance explained by sample to sample variation. Figure 9 shows all genes that were significantly associated with the fraction of mRNA from virus at a false discovery rate of 0.1. We performed the gene set analysis using the P -values from the Monocle differentialGeneTest with piano (*Väremo et al., 2013*) using the hallmark gene set from GSEA v6 (*Subramanian et al., 2005*) and Fisher's method.

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580 the Illumina deep sequencing.

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Supplementary file 1. Computer code for the analyses. This ZIP file contains a Jupyter notebook that runs CellRanger to align and annotate the reads, and a Jupyter notebook that uses Monocle to analyze the cell-gene matrix. The ZIP file also includes associated custom scripts.

Supplementary file 2. The annotated cell-gene matrix in Matrix Market Format. *[This file is too large for the eLife submission system. We will communicate with the editors to get it uploaded for a final accepted version, or we will post it on DataDryad. If you are a reviewer and need access, please contact the editor.]*

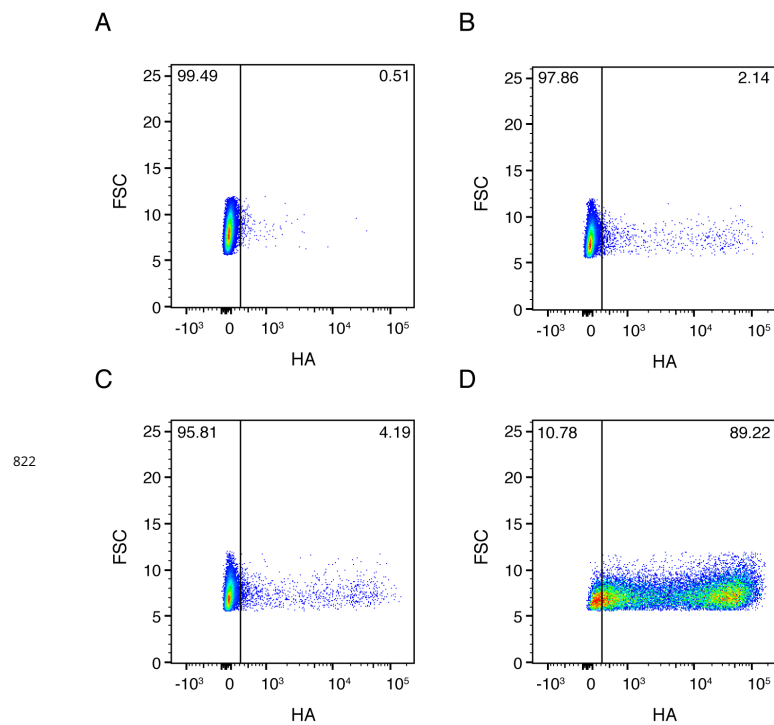


Figure 2–Figure supplement 1. Full flow cytometry data for Figure 2B. A549 cells were infected at an MOI of 0.1 as calculated by TCID₅₀ on MDCK-SIAT1 cells. **(A)** Uninfected gating control. **(B)** Cells infected with the wild-type virus stock used in our experiments. **(C)** Cells infected with synonymously barcoded virus stock used in our experiments. **(D)** Cells infected with a stock of wild-type virus propagated at a high MOI, and therefore enriched in defective particles.

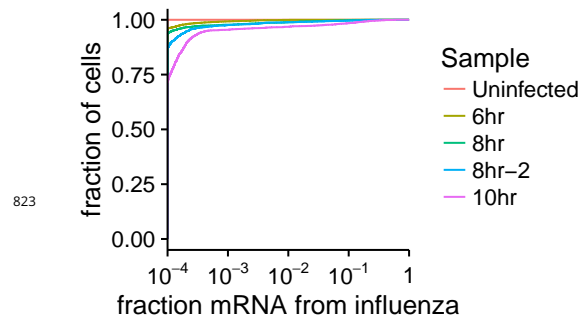


Figure 3–Figure supplement 1. For each sample, this plot shows the fraction of all cells that derive at least the indication fraction of their mRNA from influenza virus.

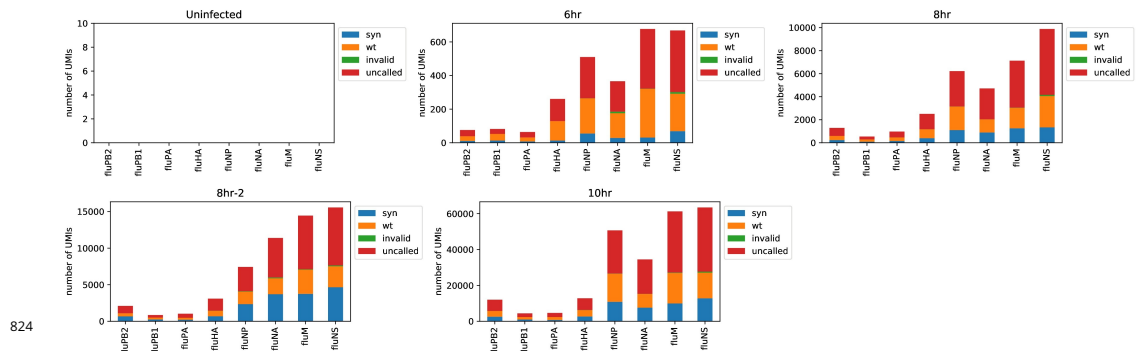


Figure 4-Figure supplement 1. The number of viral barcodes called for each sample and gene segment. Viral transcripts are classified as *syn* if they mapped to a synonymously barcoded influenza transcript, *wt* if they mapped to a wild-type influenza transcript, *invalid* if multiple reads for the same UMI differed on the status of the viral barcode, and as *uncalled* if none of the reads for that UMI overlapped the region of the viral transcript containing the viral barcode. For calculations of the number of reads in a cell derived from each viral barcode for each viral gene, the total number of detected molecules of that gene are multiplied by the fraction of those molecules with assignable barcodes that are assigned to that barcode.

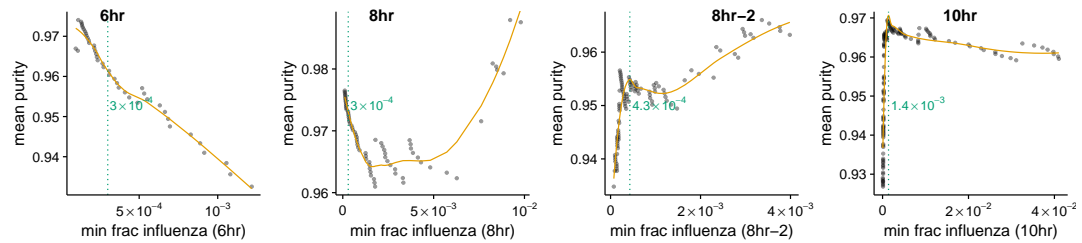


Figure 4-Figure supplement 2. Cell lysis can lead cells to the spurious association of small amounts of extraneous mRNA with individual cells. We wanted to avoid classifying as infected cells that had simply acquired such lysis-derived viral mRNA. The amount of lysis-derived viral mRNA will vary among samples as a function of both the lysis rate during the cell preparation (which always varies slightly from sample to sample in the 10X procedure) and with the amount of total viral mRNA for that sample (the more viral mRNA, the more there is to be acquired from lysed cells). As is shown in Figure 4B, the 8hr-2 and 10hr sample clearly have an enrichment of mixed barcodes in cells with small numbers of viral mRNA. For each sample, we calculated the mean purity of all cells with at least the indicated amount of viral mRNA, and determined the threshold amount of viral mRNA where purity no longer increased by finding the first maxima in a loess curve fit (orange line). We called the threshold at this point of maximum purity (dotted green line). For the 6hr and 8hr samples there is no indication of contamination from lysis-derived reads, as Figure 4B shows no increase in mixed barcodes in low viral mRNA cells. Therefore, for these samples we simply set a threshold of requiring at least 2×10^{-4} of the total mRNA to come from virus, which corresponds to ~ 2 viral mRNAs for the typical cell with 10^4 total reads (Figure 3B).

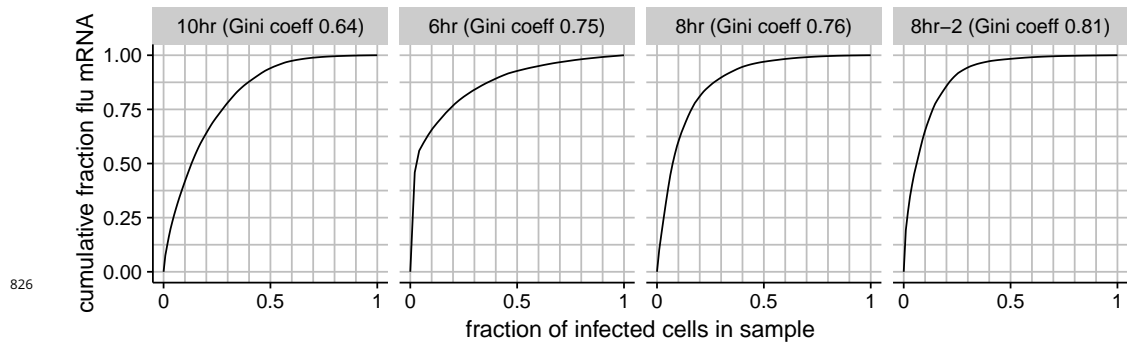


Figure 4-Figure supplement 3. The total fraction of all viral mRNA among infected cells that is attributable to a given fraction of these cells. For instance, the plot for the 8hrs sample shows that ~50% of all viral mRNA is derived from ~8% of the infected cells. The facet titles above each plot also give the Gini coefficient ([Gini, 1921](#)) that calculates the heterogeneity in the distribution of viral mRNA among infected cells. Gini coefficients of 0 indicate a perfectly even distribution across cells, and Gini coefficients of 1 indicate a maximally skewed distribution.

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[Alistair, can you add add figure]

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Figure 4-Figure supplement 4. [Alistair, can you add caption]

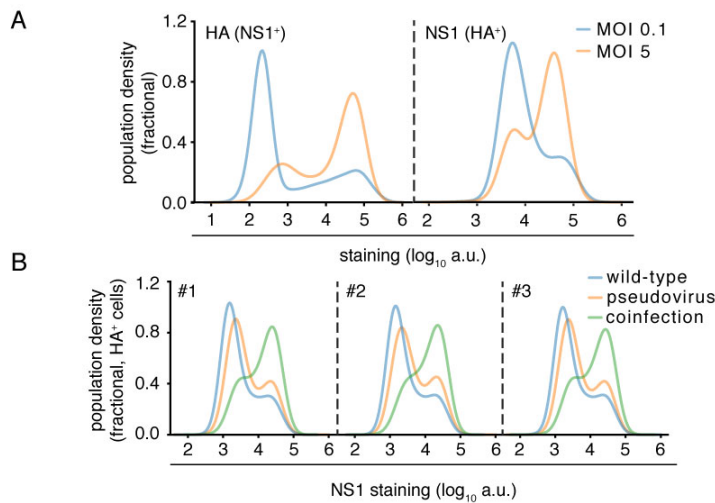


Figure 4-Figure supplement 5. (A) Flow cytometry analysis of expression of viral proteins in cells infected at high (MOI 5 as calculated by TCID₅₀ on MDCK-SIAT1 cells) or low (MOI 0.1) initial infectious dose. Cells were concurrently stained for HA and NS1 proteins 10 hours post infection. HA staining was analyzed in cells positive for NS1 (left), and NS1 staining was analyzed in cells positive for HA (right). While a higher dose leads to more cells expressing high amounts of viral protein, it does not greatly increase the amount of viral protein in either the low-expressing or high-expressing cells. Therefore, higher viral dose does not lead to a large continuous increase in viral protein production among all cells – rather, it mostly changes the proportions of cells that fall in different parts of the highly heterogeneous distribution. **(B)** Cells were co-infected with a mix of wild-type virus and pseudovirus in which the HA gene was replaced by GFP flanked by the terminal regions of the HA gene segment at an MOI of 0.1 for each virus. At 10 hours post-infection, cells were stained for NS1 and HA expression and analyzed by flow cytometry for these proteins and GFP. Cells could be annotated as infected by virions of the same type (wild-type infection indicated by presence of HA, or pseudovirus infection indicated by the presence of GFP) or both types of virions (indicated by presence of HA and GFP). Coinfected cells, like cells infected at a higher infectious dose, occupy different positions in the distribution of viral protein production but do not exhibit a continuous increase in viral protein production. Replicates shown.

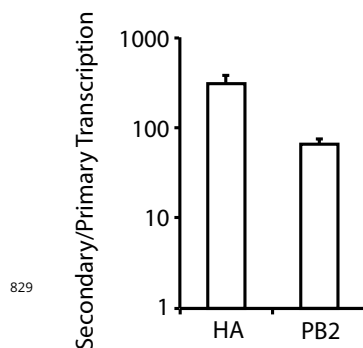


Figure 5-Figure supplement 1. A549 cells were infected at an MOI of 0.2 as calculated on MDCK-SIAT1 cells in either the presence or absence of the protein-translation inhibitor cyclohexamide, and viral mRNA was quantified at 8 hours post-infection by qPCR. The cyclohexamide prevents translation of new PB2, PB1, PA, and NP protein, and so prevents the formation of the new RNPs needed for secondary transcription. The bars show the relative amount of HA and PB2 mRNA in the absence versus the presence of cyclohexamide. Error \pm S.D. n=3.

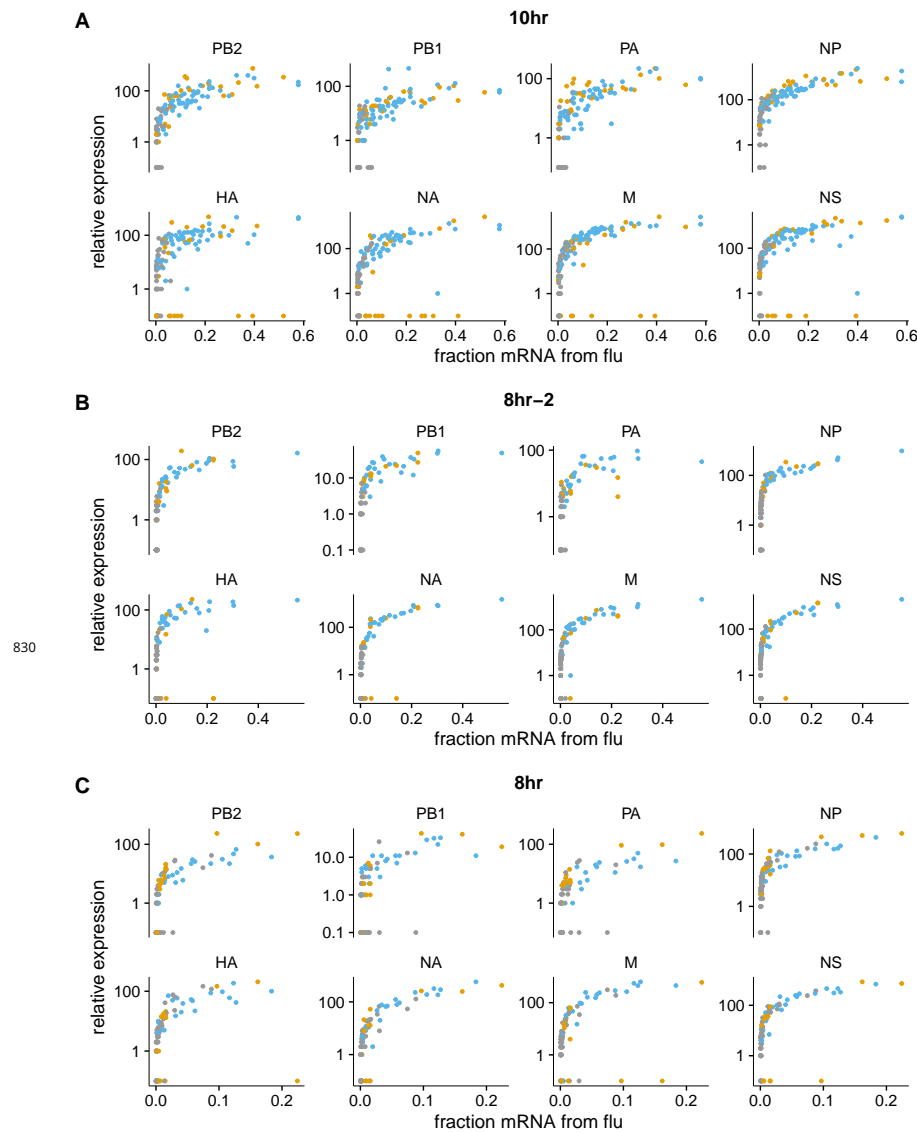


Figure 5–Figure supplement 2. The normalized expression of each viral gene as a function of the fraction of total mRNA derived from virus, shown for the 10-hour and 8-hour samples individually (the other samples had too few infected cells for this analysis to be useful). Points are colored as in Figure 5A.

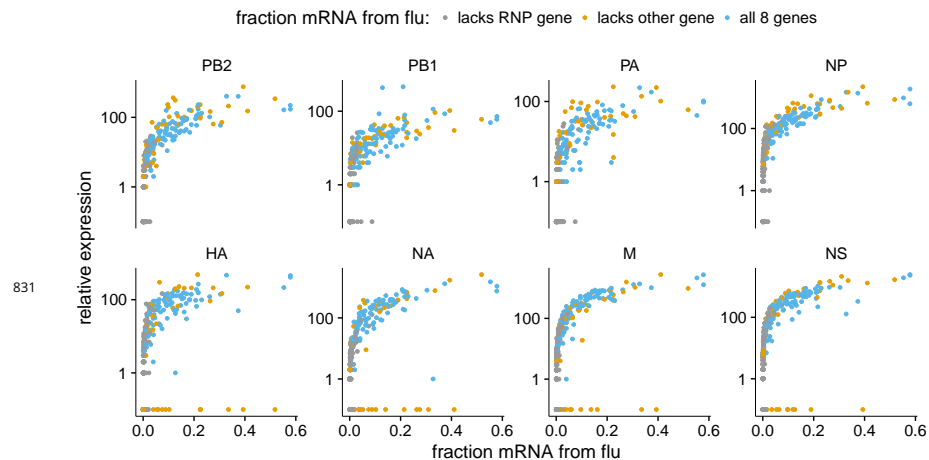


Figure 5-Figure supplement 3. The normalized expression of each viral gene as a function of the fraction of total mRNA derived from virus, excluding cells that were annotated as coinfected based on the presence of both viral barcodes.

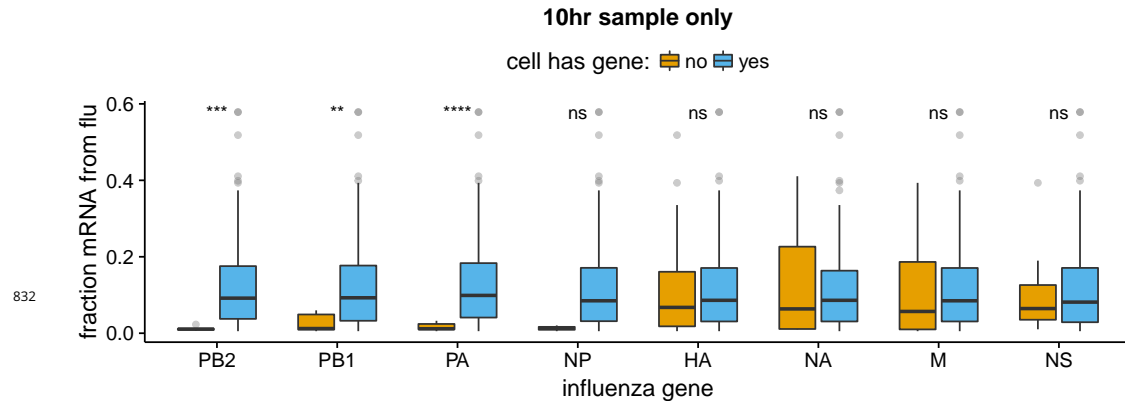


Figure 5-Figure supplement 4. The absence of viral RNP genes but *not* non-RNP genes remains significantly associated with reduced viral burden when we examine only the 10-hr sample, which is the single time point with the most data points. The difference for NP is no longer statistically significant due to low counts of infected cells lacking NP, but the trend remains. We do not show statistical analyses for other samples, as the number of infected cells is too low.

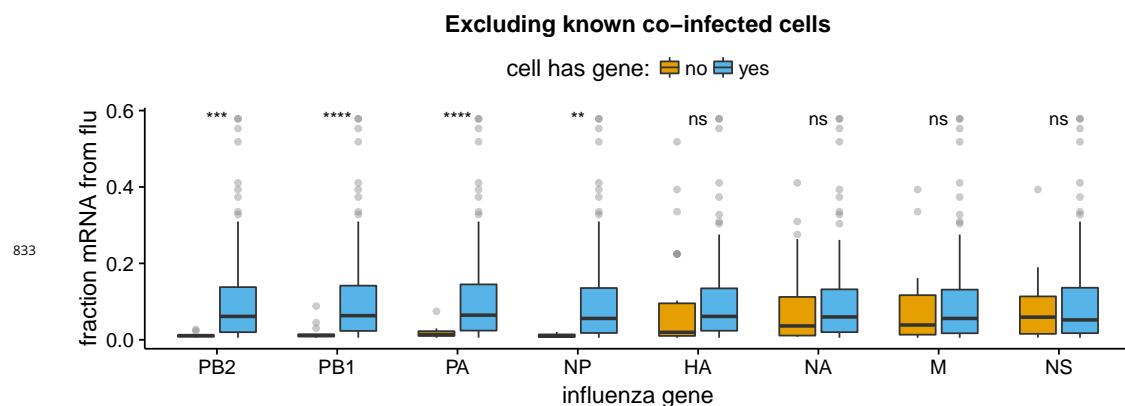


Figure 5-Figure supplement 5. All findings in Figure 5B remain unchanged if we exclude cells called as coinfecte based on the presence of mixed viral barcodes.

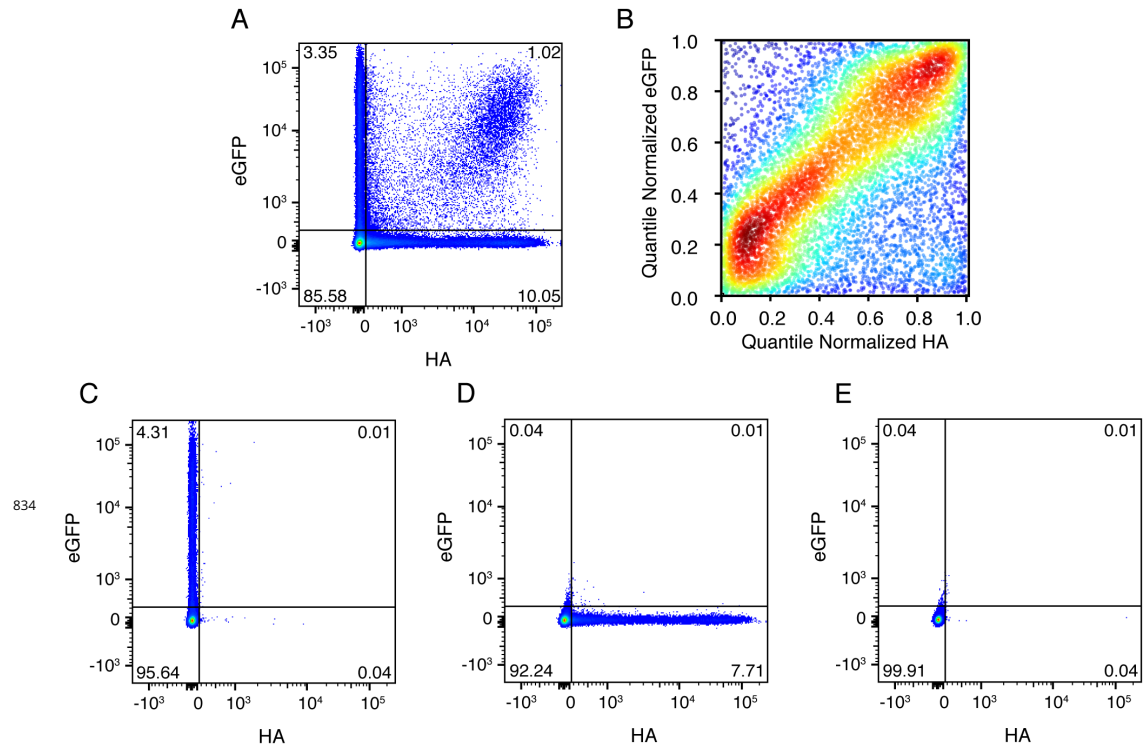


Figure 7-Figure supplement 1. **(A)** Cells were co-infected with a mix of wild-type virus and virus in which the HA gene was replaced by GFP flanked by the terminal regions of the HA gene segment. At 10 hours post-infection, cells were analyzed by flow cytometry for HA and eGFP expression. **(B)** The expression of HA and GFP are correlated in co-infected cells. Shown are the quantile-normalized HA and eGFP signals for double-positive cells. Cells are colored by density, using a Gaussian kernel density estimate. **(C),(D),(E)** Gating controls, single infection with eGFP virus, single infection with wild-type virus, and uninfected cells, respectively.

Category	Genes
Detoxification	AKR1C3, AKR1B10, GPX2, ALDH3A1, ALDH1A1, NQO1, CBR1, PRDX1
Protein folding	TXN, PPIA
Electron transport chain	NDUFB4, MT-CO1, MT-CO3
Regulators	ATF3, GADD45B
ROS-responsive relevance complex/unknown	UBB, NME1

Figure 9-Figure supplement 1. Table delineating genes in Figure 9 that are associated with the response to oxidative stress (*Duong et al., 2017; Jung et al., 2017; Lee and Ryu, 2017; Peuchant et al., 2017; MacLeod et al., 2016; Jiang et al., 2016; Gorrini et al., 2013; Miura et al., 2013; Kim et al., 2009; Banning et al., 2005; Murray et al., 2003; Doyle et al., 1999*).

[add figure]

Figure 9–Figure supplement 2. [Alistair, can you add figure and caption showing the increase in the percent of flu positive cells after treatment. It should just be possible to put the methods for this experiment in the figure caption.]

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