Transcriptional dynamics of influenza virus infection at the single-cell level

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- Abstract Influenza virus infection induces large changes in cellular transcription. Previously this has mostly been looked at using bulk measurements Here we examine the process at the level of single cells. We find extremely wide variation in the extent of viral gene transcription across infected cells. IFN induction is very rare. Some cellular pathways may be consistently altered in cells with high burden of viral transcripts. Overall, highlights remarkable heterogeneity in the outcome of infection.

Introduction

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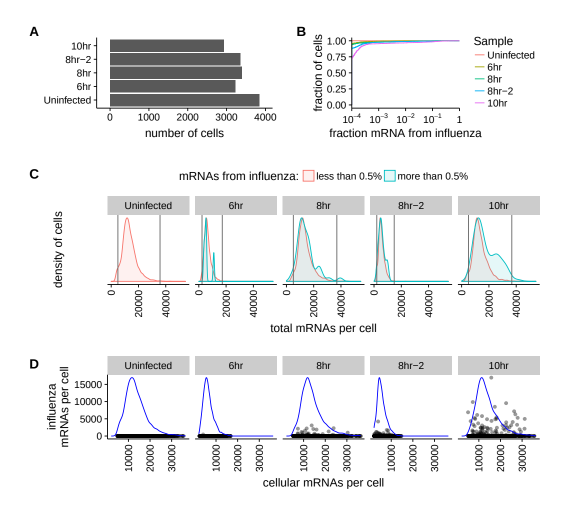


Figure 1. Overview of amounts of cellular and influenza virus mRNAs detected in each cell. **(A)** Number of cells captured for each sample. **(B)** Cumulative fraction plot showing the amount of mRNA derived from influenza for each sample. In all samples, most cells had little or no influenza mRNA. **(C)** Distribution of number of cellular and viral mRNAs per cell in each sample. Cells outside the vertical gray lines were considered outliers and excluded from subsequent analyses. **(D)** The number of cellular and viral mRNAs for each cell is plotted as a point. The blue lines show the overall distribution of the number of cellular mRNAs per sample. At later timepoints, a small number of cells had a very high number of viral mRNAs.

Figure 1-Figure supplement 1. Shorter caption for main text.

Figure 1-Figure supplement 2. This is another supplementary figure.

Figure 1-source data 1. This is a description of a data source.

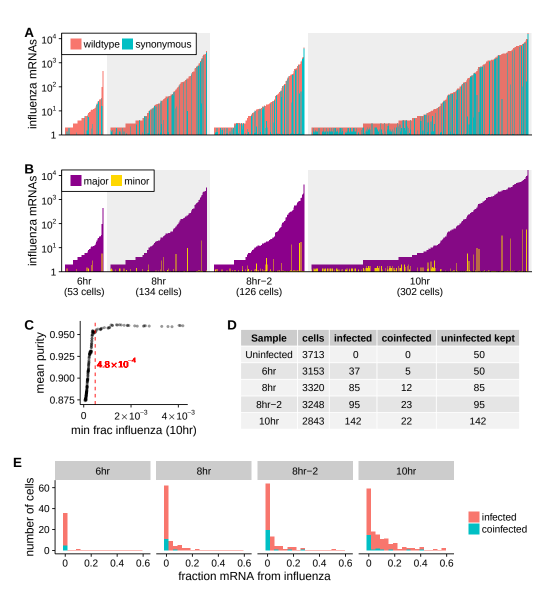


Figure 2. Synonymous barcodes near the 3' end of the influenza virus mRNAs were used to identify co-infection and distinguish true infections from cells that contained a few spurious viral reads. (A) For all cells with at least two viral mRNAs for which the synonymous barcode could be called, each line is proportional to the logarithm of the number of viral mRNAs in that cell. The bars are colored in linear proportion to the fraction of the viral mRNAs derived from either wildtype or synonymously barcoded virus. (B) Same as (A), but now each bar is colored according to the relative proportions of the more common (major) and less common (minor) barcoded virus variant. At low levels of viral mRNA there is often a roughly equal mix of barcodes, since many of these cells have simply picked up environment mRNA which is about equally likely to derive from either virus. But at higher levels of viral mRNA, truly infected cells are mostly one pure barcode except for a few cells that are truly co-infected. (C) We determined a cutoff for calling "true" infections by fitting a curve to the mean barcode purity of all cells with greater than a given fraction of their mRNA derived from virus. We called the cutoff at the point at which purity stops increasing with the fraction of viral mRNA. (D) The number of cells identified as infected and co-infected for each sample. For all samples, the vast majority of cells were not infected, so for subsequent analyses we subsampled to a number of uninfected cells that was the greater of 50 or the number of infected cells. (E) The distribution of the fraction of mRNA derived from virus for each sample for both infected and co-infected cells. For all samples, there is a very wide distribution of the amount of viral mRNA.

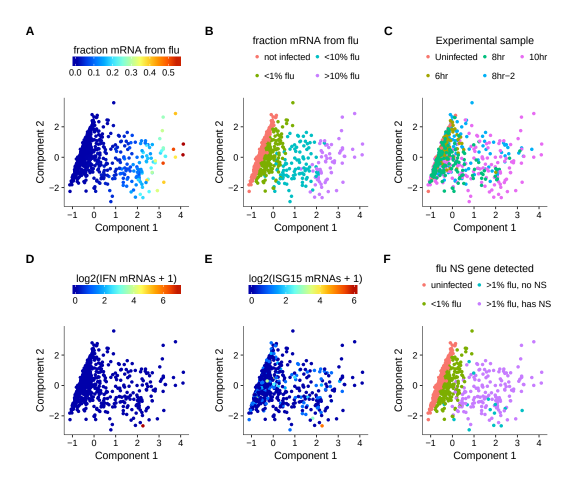
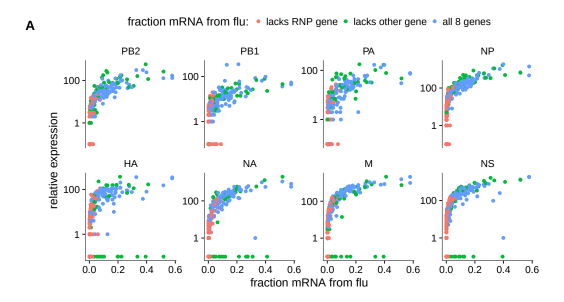


Figure 3. Visual layout of the cells according to "pseudotime". The layout is the same in all panels, but each panel colors the cells according to a different property. **(A), (B)** Cells colored by the fraction of their mRNA that is viral. **(C)** Cells colored by experimental sample. While it is clear that cells from later timepoints often have more viral RNA, there are cells from earlier timepoints with a high viral burden and cells from late timepoints with a low viral burden. **(D)** Cells colored by the number of type I and III interferon transcripts detected. Only one cell has high expression of these interferons. **(E)** Cells colored by the expression of the interferon-stimulate gene ISG15. **(F)** For cells with at least 1% of their reads from influenza, are the cells expressing the viral NS protein? The one interferon-positive cell is lacking NS, but many other cells also lack NS but do not express interferon.



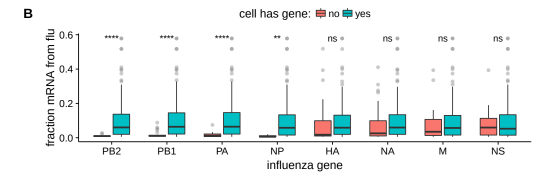


Figure 4. The viral infection burden in individual cells as a function of the amount of each viral gene detected. **(A)** Fraction of mRNAs in each cell derived from virus as a function of the *normalized* expression of each viral gene in that cell. This plot shows that all cells with very high viral burden express all of the RNP genes, but some cells with high viral burden lack each of the other four viral genes. **(B)** Statistical tests confirming that absence of viral RNP genes is significantly associated with reduced viral burden, but that the absence of the non-RNP genes does not lead to a clear decrease in viral burden.

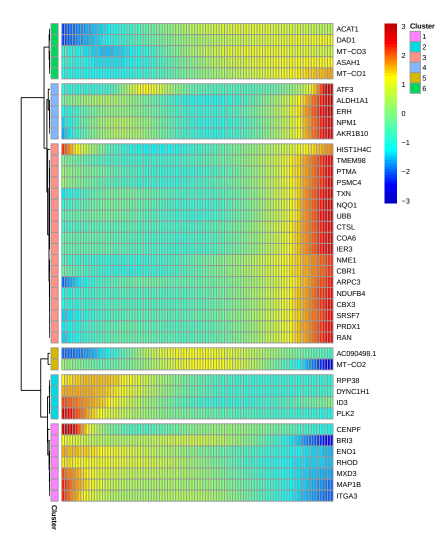


Figure 5. Cellular genes that are differentially expressed with respect to the amount of influenza mRNA in individual cells infected with full influenza virus containing all eight genes. Shown are all genes differentially expressed with Q < 0.1.

Figure 5-source data 1. The full results of the differential expression test is in p_sig_cellular_genes.csv. **Figure 5-source data 2.** The results of a gene-set analysis are in p_sig_cellular_genes.csv.

38 Discussion

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48 Methods and Materials

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58 Some LATEX Examples

Use section and subsection commands to organize your document. LeteX handles all the formatting and numbering automatically. Use ref and label commands for cross-references.

61 Figures and Tables

62 If you use the following prefixes for your \label:

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Figures fig:, e.g. \label{fig:view}
Tables tab:, e.g. \label{tab:example}
Equations eq:, e.g. \label{eq:CLT}
Boxes box:, e.g. \label{box:simple}
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you can then use the convenience commands as in \FIG{cells}, to generate cross-reference??.

68 Citations

LaTeX formats citations and references automatically using the bibliography records in your .bib
file, which you can edit via the project menu. Use the \cite command for an inline citation, like
Trapnell et al. (2014), and the \citep command for a citation in parentheses (Trapnell et al., 2014).
The LaTeX template uses a slightly-modified Vancouver bibliography style. If your manuscript is
accepted, the eLife production team will re-format the references into the final published form. It is

not necessary to attempt to format the reference list yourself to mirror the final published form.

Acknowledgments

Additional information can be given in the template, such as to not include funder information in the acknowledgments section.

References

Trapnell C, Cacchiarelli D, Grimsby J, Pokharel P, Li S, Morse M, Lennon NJ, Livak KJ, Mikkelsen TS, Rinn JL.
 Pseudo-temporal ordering of individual cells reveals dynamics and regulators of cell fate decisions. Nature
 Biotechnology. 2014; 32(4):381.



Figure 1–Figure supplement 1. This is a supplementary figure's full caption, which will be used at the end of the manuscript.



Figure 1-Figure supplement 2. This is another supplementary figure.