22Feb19: Neutralization assays against WT Perth/2009 + WSN NA GFP virus with Vietnam serum samples

"A-H" will be referred to as columns, and "1-12" as rows. I had the plate with column A-H going left to right, and rows 12-1 from top to bottom.

| | COLONN | | | | | | | |
|-----|----------|------------|--------------------|----------------|----------------|----------------|--------------------|----------|
| ROW | Α | В | С | D | E | F | G | Н |
| 1 | NAM only | virus only | virus + cells only | serum dilution | serum dilution | serum dilution | virus + cells only | NAM only |

I used the "Perth/2009 HA + WSN NA GFP virus" that Juhye rescued on 20180905. Based on the MOI test Juhye conducted, I will use an MOI of 0.5

| Serum Sample | Vol of Ab for serial dil | Time start virus + Ab | Time start virus+Ab+cells |
|---------------|--------------------------|-----------------------|---------------------------|
| 16 (HC080048) | 40 uL (3-fold dilutions) | 2:07 | 3:44 |
| 17 (HC080043) | 40 uL (3-fold dilutions) | 2:09 | 3:47 |
| 18 (HC060106) | 40 uL (3-fold dilutions) | 2:11 | 3:50 |
| 19 (HC140010) | 40 uL (3-fold dilutions) | 2:13 | 3:52 |
| 20 (HC070072) | 40 uL (3-fold dilutions) | 2:15 | 3:55 |
| 21 (HC070041) | 40 uL (3-fold dilutions) | 2:17 | 3:57 |
| 22 (HC120043) | 40 uL (3-fold dilutions) | 2:19 | 4:00 |
| 23 (HC150036) | 40 uL (3-fold dilutions) | 2:20 | 4:03 |

For multichannel, used Fisher brand blue box/green wafer tips, #02-707-431

- 1. Thawed sera and made dilutions to working stock solutions using NAM
 - Diluted 16.67 uL of serum in 133.33 ul NAM for 150 uL total volume
- 2. Added 80 uL NAM to all wells of all 8 plates (~65 mL of NAM). Also began thawing viruses
- 3. Added 40 uL of the diluted serum to the top row (row 12) of the plate for columns D, E, and F. Did this for each plate
- 4. Using the multichannel with three tips attached, mixed the top row D/E/F and transferred 40 uL the second row, mixed, etc. Removed 40 uL from the last row. Did not change tips in between rows (not necessary here)
- 5. Added 40 uL NAM to column B of all plates (virus only column) to make up for no cells being added to these columns.
- 6. Prepared virus inoculum. Diluted Perth/09 HA + WSN NA GFP 48 hpt virus to 500 IP/uL. (~3.5 mL per plate= 28 mL total)
 - Titer = 2419 IP/uL. Make MM of virus (5.79 mL virus + 22.2 mL NAM) need 6 aliquots of virus
- 7. Using a reservoir and multichannel added 40 uL of virus to columns B-G. Used new tips for each row, but did not mix virus and Ab (no need to mix). Wrote down time once virus was added to all wells of a plate, and incubated at 37°C.
- 8. While virus + antibody mixtures were incubating, started preparing cells by trypsinizing MDCK-SIAT1-PB1-TMPRSS2 cells, neutralizing with D10, spinning at 1200 rpm for 4 min, then resuspending in 20 mL NAM
- 9. Made a master mix of cells at 1e6 cells/mL, making a 28 mL master mix (~3.5 mL per plate)
 - Counted <u>2.29e6</u> cells/mL =>12.23 mL cell culture + 15.77 mL NAM
- 10. After ~1.5 hr of virus + Ab incubation, took each plate out of the incubator in order, and added 40 uL cells to columns C-G, using fresh tips for each row.

 Used a fresh reservoir for each plate, re-mixed cell suspension before adding to reservoir. Wrote down time cells were added to the wells of a plate

 11. At ~18 hpi (9:30 AM), read on plate reader with "Neutralization assay 96 well" program.