

## Station B: Chemagic Viral 300 DNA/RNA Kit

*This protocol starts immediately after lysing the sample and starts at step 2 with the addition of magnetic beads and binding buffer 2*

### Code parameters:

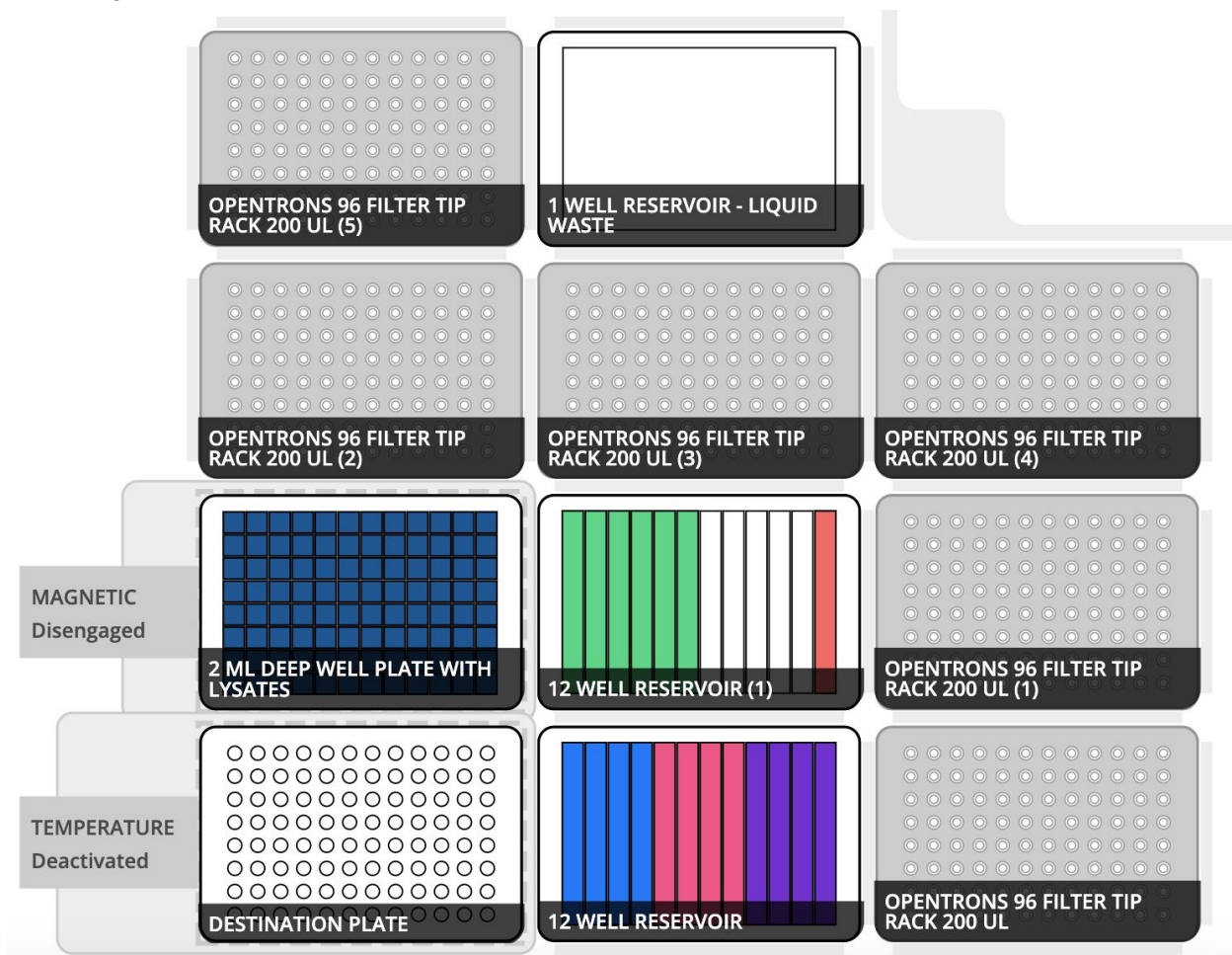
- Change the sample number on line 14 (default is 8, max is 94)
- Change the elution volume on line 15 (default is 50µl, max is 100µl)
- Change the sample starting volume on line 16 (default is 420µl)
- Tip rack tracking can be changed from False to True on line 17 (default is False)
- Make “tiprack parking” True or False on line 18\* (default is True)

*\*if you selected True for “tiprack parking,” tips used for the same buffers with the same samples will be reused where 1 tiprack turns into a tiprack where used tips are “parked”. This method has low risk of contamination and is highly recommended to avoid pauses to refill tipracks.*

### Pipettes:

- P300 multichannel on the left mount

### Deck Layout:



**Labware and module requirements:**

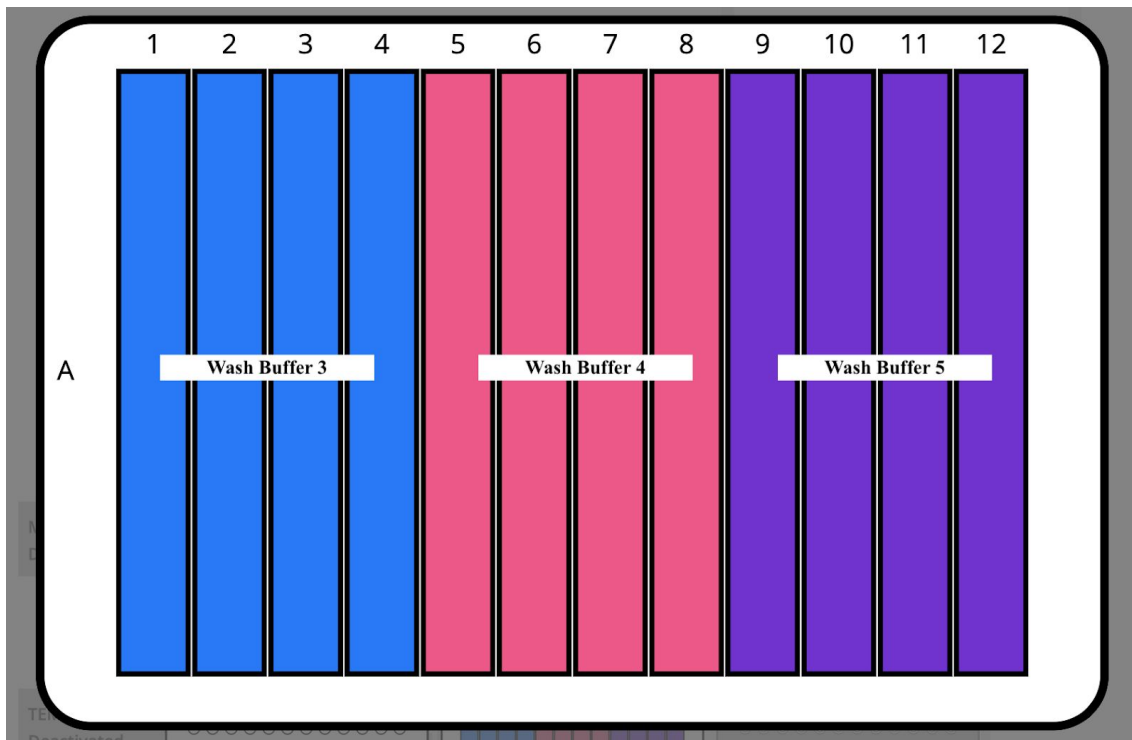
- 1 x magnetic module
- 1 x temperature module
- 1 x 2mL deep well plate **[input with lysates]**
- 6 x 200µl filter tip racks (10 x 200µl if you select false for tiprack parking)
- 1 x 1 well reservoir **[loaded empty for the liquid waste in slot 11]**
- 2 x 12 well reservoir with reagents **[1 x holds Magnetic Beads & Binding Buffer 2 Mixture, and Elution Buffer 6, 1 x holds Wash Buffer 3, Wash Buffer 4, and Wash Buffer 5]**
- 1 x 96 well aluminum block loaded on top of the temperature module in slot 1
- 1 x 96 well PCR plate **or** PCR strip tubes to match the number of samples **[output with eluates/extractions]**

**Volume requirements:**

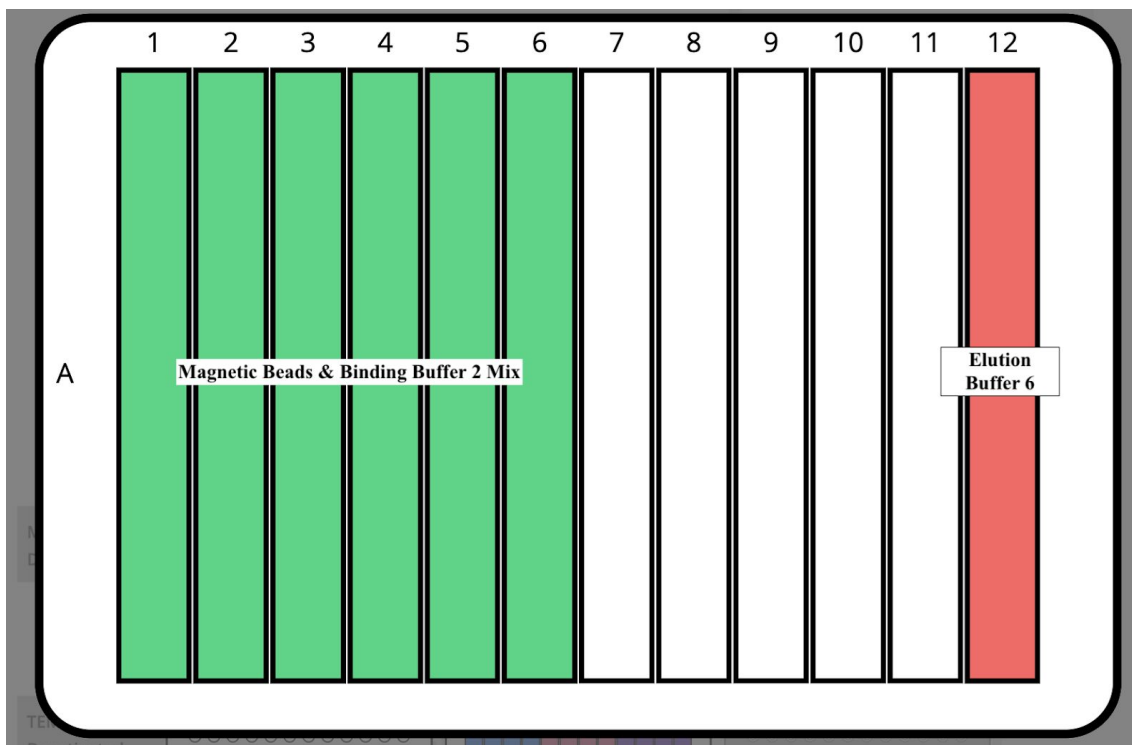
**Note:** the below volumes account for a dead volume - the dead volume can be adjusted depending on the calibration of the pipette to the labware, but it's best to have a dead volume of at least 10%

Reagents	Volume per sample (µl)	Volume for 8 samples (µl)	Volume for 48 samples (mL)	Volume for 96 samples (mL)
<b>Magnetic Beads &amp; Binding Buffer 2</b>	620	7,440	32.2	62
<b>Wash Buffer 3</b>	500	6,000	26	50
<b>Wash Buffer 4</b>	500	6,000	26	50
<b>Wash Buffer 5</b>	500	6,000	26	50
<b>Elution Buffer 6</b>	50 - 100	600 - 1,200	2.6 - 5.2	5 - 10

### Slot 2 - 12 well reservoir setup



### Slot 5 - 12 well reservoir setup



### Before you begin:

1. Pre-cool the Temperature Module in the Opentrons App to 4°C
2. Create the **Magnetic Beads & Binding Buffer 2 Mixture**

	Volume per sample	Volume for 8 samples	Volume for 48 samples	Volume for 96 samples
<b>Magnetic Beads</b>	20 µl	240 µl	1,040 µl	2 mL
<b>Binding Buffer 2</b>	600 µl	7,200 µl	31.6 mL	60 mL

**Note:** the beads settle quickly so be sure to vortex the solution thoroughly before adding the mixture to the reservoir

3. Add the Wash Buffer 3, Wash Buffer 4, and Wash Buffer 5 to the 12 well reservoir in slot 2
4. Add the Magnetic Bead & Binding Buffer Mixture and Elution Buffer 6 to the 12 well reservoir in slot 5
5. Place the deep well plate filled with lysates on top of the magnetic module in slot 4.
6. Add a 96 well aluminum block and the 96 well PCR plate or PCR strip tubes on top of the Temperature Module in slot 1

The final plate of eluates/extractions will be found on top of the temperature module in slot 1. Once the run is complete, please proceed to Station C for RT-PCR set up.