# 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:**

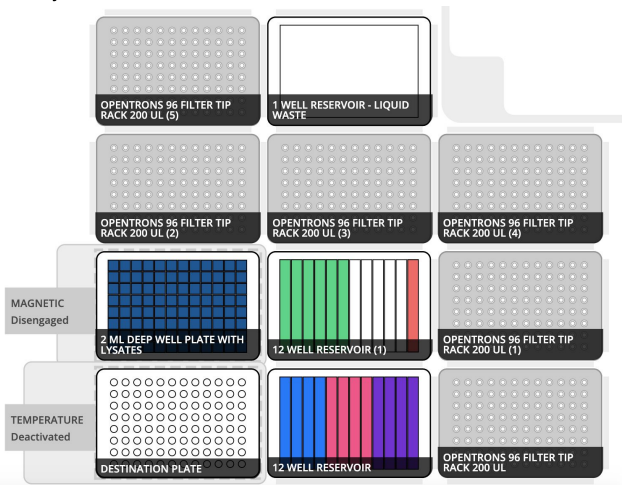
- Change the sample number on line 14 (default is 8, max is 94)
- Change the elution volume on line 15 (default is 50µl, max is 100µl)
- Change the sample starting volume on line 16 (default is 420µl)
- Tip rack tracking can be changed from False to True on line 17 (default is False)
- Make "tiprack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

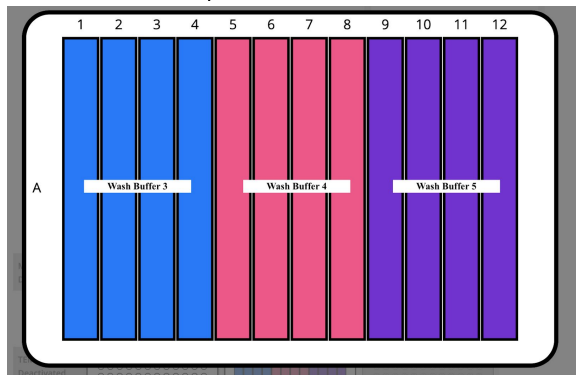
- 1 x magnetic module
- 1 x temperature module
- 1 x 2mL deep well plate [input with lysates]
- 6 x 200µl filter tip racks (10 x 200µl if you select false for tiprack parking)
- 1 x 1 well reservoir [loaded empty for the liquid waste in slot 11]
- 2 x 12 well reservoir with reagents [1 x holds Magnetic 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:**

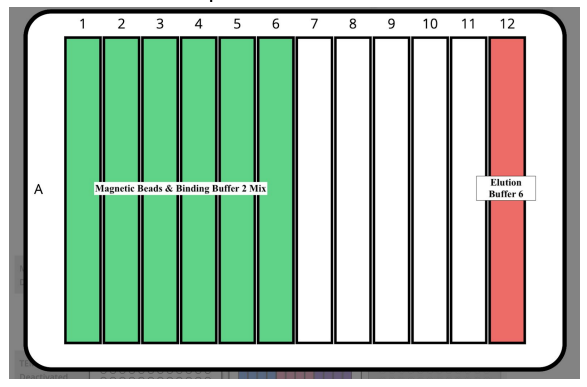
**Note**: the below volumes account for a dead volume - the dead volume can be adjusted depending on the calibration of the pipette to the labware, but it's best to have a dead volume of at least 10%

Reagents	Volume per sample (µI)	Volume for 8 samples (µI)	Volume for 48 samples (mL)	Volume for 96 samples (mL)
Magnetic Beads & Binding Buffer 2	620	7,440	32.2	62
Wash Buffer 3	500	6,000	26	50
Wash Buffer 4	500	6,000	26	50
Wash Buffer 5	500	6,000	26	50
Elution Buffer 6	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
Magnetic Beads	20 μΙ	240 μΙ	1,040 μΙ	2 mL
Binding Buffer 2	600 µI	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.