SARS-CoV-2 Antigen Detection ELISA V.1

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XPRIZE Rapid Covid Testing

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Protocol status: Working

Created: Sep 07, 2020

MATERIALS

- Tween 20 Sigma Aldrich Catalog #P7949
- Phosphate Buffered Saline (25x) pH 7.6 ScyTek Catalog #PBS-20000
- Sodium Bicarbonate Sigma Aldrich Catalog #S6297
- Hydrochloric Acid Fischer Scientific Catalog #A142-212
- BSA Sigma A3059 Sigma Aldrich Catalog #A3059
- Blocker™ Casein in PBS Thermo Fisher Scientific #37528 Thermo Fisher Scientific Catalog #37528
- ELAST® ELISA Amplification System Perkin Elmer #NEP116E001EA Perkin Elmer Catalog #NEP116E001EA
- HRP Conjugation Kit - Lightning-Link Abcam #ab102890 Abcam Catalog #ab102890
- SureBlue Reserve TMB 1-Component Microwell Peroxidase Substrate KPL Cat #53-00-03 KPL Catalog #53-00-03
- Deionized (DI) Water (dan use lab DI water) Contributed by users
- Purified recombinant capture and detection antibodies (see Table under Materials Text) Contributed by users

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<th>Antibody</th>
<th>Binding domain*</th>
<th>Species</th>
<th>Isotype</th>
<th>Description</th>
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<td>DH1219</td>
<td>NP</td>
<td>Human</td>
<td>IgG1</td>
<td>NP capture</td>
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</tbody>
</table>
**Keywords:** SARS-CoV-2

**Reagents**

1. Tween 20, Sigma Cat. No. P7949
2. Phosphate Buffered Saline (25x) pH 7.6 from ScyTek Laboratories, Cat No: PBS-20000
3. Sodium Bicarbonate, Sigma, Cat No: S6297
4. Hydrochloric Acid, Fischer Scientific Cat No. A142-212
5. BSA Sigma #A3059
6. Blocker™ Casein in PBS Thermo Fisher Scientific # 37528
7. ELAST® ELISA Amplification System Perkin Elmer # NEP116E001EA
8. HRP Conjugation Kit - Lightning-Link Abcam # ab102890
9. SureBlue Reserve TMB 1-Component Microwell Peroxidase Substrate, KPL Cat No. 53-00-03
11. Purified recombinant capture and detection antibodies:

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<td>DH1219</td>
<td>NP</td>
<td>Human</td>
<td>IgG1</td>
<td>NP capture</td>
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</table>

*SARS-CoV-2 Spike (S) and Nucleocapsid/nucleoprotein (NP)*

**Materials and Equipment**

1. Costar 3700 high binding 384 well plates
2. Plate Washer: Biotek 406
3. Plate Reader: Molecular Devices 384plus
**Preparation of Storable Solutions**

### 0.1 M Sodium Bicarbonate (Coating buffer)

1. Measure 8.4 g of Sodium Bicarbonate. Transfer to a 1L bottle.
2. Bring the volume up to 1 L using DI water. Mix solution until dissolved.
3. Filter the 0.1M Sodium Bicarbonate solution using Corning® 500mL Vacuum Filter/Storage Bottle System with 0.22µm Filter Membrane in order to remove any undissolved particles.
4. Transfer the solution into new bottle, and store at room temperature. Discard after one month.

### 1%BSA/0.1%Tween-20 in PBS (1%BSA/T)

1. 10 g of BSA. Transfer to 1L bottle.
2. Add 40 mLs of 25X PBS and 1.0 mL of Tween 20. Bring volume to 1L with DI water. Stir the solution until dissolved. Store at 4°C. This has a shelf life of two weeks. If any turbidity occurs, discard and remake.

### Assay Wash (10L)

1. In 10L bottle, add 10 mL of Tween20 and 400 mL of 25X PBS. Bring the volume up to 10L using DI water. Stir completely. This will yield in 1X PBS/0.1%Tween-20 solution.

### Acid Stopping Solution (1X HCl Solution)

1. To make 1L of 1X HCl (.33 N) solution using recommended HCl, add approximately 700 mL of DI water into 1L bottle.
2. Add 27 mL of Hydrochloric Acid. Bring the volume to 1L with DI water. Mix the solution.

**Assay Procedure**

4. Dilute each capture antibody, DH1041 (S) and DH1219 (NP), to a final concentration of 4ug/ml in coating buffer at a volume sufficient to fill all assay wells of a 384 well plate with 15ul (see plate map at the end). Unused portions of the plate can be left empty if desired.
5 Coat assay wells of high-binding 384 well ELISA plate with 15 µL/well of coating solution. Tap plate(s) lightly to ensure all liquid settles at the bottom of the wells. Cover tightly or wrap the plate(s) with plastic wrap then incubate overnight at 4°.

6 Wash plate(s) twice with 100µL of assay wash per well. Make sure wells are empty (pat plates on paper towel if necessary).

7 Block wells with 80µL casein/well. Tap plates lightly to ensure all liquid settles at the bottom of the wells. Incubate for at least 2 hours at room temperature or overnight at 4oC. If storing overnight, wrap the plate(s) with plastic wrap.

8 Purified Recombinant Antigen Standard curve:
   - In separate tubes dilute the recombinant S and NP proteins to 1ug/ml in 1%BAS/T.
   - In a 96 well dilution plate perform a serial dilution in three-fold dilution ratio. Have the first well start at 0.01ug/ml (10,000pg/pm) for each protein.
   - This requires a 1 to 100 dilution from each individual protein tube (at 1ug/ml) into the same starting well. Example: In the first well add 265ul 1%BAS/T. Then add 2.5ul of each 1ug/ml tube of S and NP.
   - Into all other wells add 180ul of 1%BAS/T. Then transfer 90ul from the first well into the next well, serially through all 12 wells, with proper mixing.

9 Vortex or invert the test samples to mix, then centrifuge 10,000 g x 5 minutes to settle any solids.

10 After blocking the plate(s), wash one time with assay wash. Make sure wells are empty. All further steps are conducted at room temperature.

11 Following the plate layout add 10ul/well undiluted sample in triplicate to the assay plate.
Add standard curve to the assay plate at a volume of 10ul/well in duplicate.

To blank rows add 1%BSA/T for subtraction later.

Tap plate to make sure all samples are at the well bottom. Cover and incubate for 60 minutes.

During incubation, prepare the HRP labeled detection antibody cocktail by diluting each HRP labeled antibody to its optimal concentration in the same tube 1%BSA/T. Currently DH1046 (anti-S) is used at 0.25ug/ml and DH1218 (anti-NP) is used at 0.1ug/ml.

- The HRP labeled antibodies must be pre-labeled following the procedure outlined in the following kit: HRP Conjugation Kit - Lightning-Link Abcam #ab102890.
- The working concentration of each HRP labeled antibody must be determined by titration in a prior optimization assay as outlined briefly below:
  1. Using all reagents outlined above, coat plate with the capture antibodies as specified. Then wash and block as above.
  2. Prepare recombinant antigen standards by diluting each antigen separately to a final concentration of 1,000pg/ml. Add each antigen solution separately to the assay plate as outlined above.
  3. Prepare a dilution of each HRP detection antibody separately, beginning at 1ug/ml serially diluting for 12 wells.
  4. Incubate the HRP-detection antibody for 60 min and continue the detection as outlined in this protocol.
  5. Choose a working concentration of each HRP detection antibody that is both at saturation and practical, i.e >=0.1ug/ml.

During incubation calculate and prepare the ELAST ELISA enhancer reagents according to the protocol provided in the kit specified in the reagents section.

- BT Solution: Volume required = 10ul/well x # of wells + 10%. Example: If 4ml is needed (2ml DiH2O + 2ml BT diluent + 40ul BT reagent)
- HRP Solution: Volume required = 10ul/well x # or wells + 10%. Dilute HRP 1 to 500 in 1%BST/T. Example: If 4ml is needed (1%BSA/T + 8ul HRP reagent). Protect from light until use.

Following incubation with HRP detection antibodies, wash plate(s) twice. Make sure wells are empty.
18 To all wells add 10µL/well of BT solution. Incubate plate for 20 minutes.

19 Wash plate(s) four times.

20 Add 10µL of HRP solution. Incubate plate for 20 minutes.

21 Wash plate six times.

22 Add 20µL/well of TMB substrate. Incubate 12 minutes.

23 Stop reaction with addition of 20µL acid stop solution. Tap plates to distribute stop solution.

24 Read plates at 450 nm between 5 to 30 minutes after stopping the reaction.

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**Determination of Results**

25 ■ Average all background wells.
- Subtract background average from all wells.
- Average each set of triplicates, and the standard curve duplicates.
- Any sample greater than OD 0.15 after subtraction is considered positive.
- The standard curve should be positive to at least 40pg/ml.