1. Introduction

Influenza antigen reagent TGA Lot 2017/118B is prepared for single radial immunodiffusion assay (SRID) of A/Singapore/GP1908/2015 (IVR-180A) egg-derived antigens using an appropriate TGA antiserum reagent.

Note: A/Singapore/GP1908/2015 reassortants IVR-180 and IVR-180A are derived from the same reassortant virus clone, differing by a single passage in eggs. The HA and NA sequence of the two passage levels are identical.

2. Unitage

Assigned potency of antigen Reagent Lot 2017/118B:

74 μg (microgram) of HA per mL, after reconstitution of the lyophilized reagent in 0.5 mL of distilled water, or:

37 μg (microgram) of HA per vial.

Antigen Reagent Lot 2017/118B was calibrated using sheep antiserum Lot AS418 raised against egg-derived A/Singapore/GP1908/2015 (IVR-180).

For further information please contact: influenza.reagents@health.gov.au

3. Contents

Country of origin of biological material: Australia

Antigen Reagent Lot 2017/118B was produced in non-SPF, embryonated chicken eggs and was obtained from a production zonal pool (ZP), which has been inactivated by 0.1% v/v beta-propiolactone (βPL). Sucrose was removed from the inactivated strain virus, prior to dilution with an equal volume of 6% w/v dextran in 0.9% w/v sodium chloride, mixed thoroughly and dispensed for freeze-drying in 0.5 mL volumes as described by Campbell, P.J.; Journal of Biological Standardisation, 1974, 2, 249-267.

The mean of vial weights was 0.506 g with a coefficient of variation of 0.66 %.
4. Caution

**THIS PREPARATION IS NOT FOR ADMINISTRATION TO HUMANS.**

The preparation does not contain material of human origin.

As with all materials of biological origin, this preparation should be regarded as potentially hazardous to health. It should be used and discarded according to your own laboratory's safety procedures. Such safety procedures probably will include the wearing of protective gloves and avoiding the generation of aerosols. Care should be exercised in opening ampoules or vials, to avoid cuts.

5. Use of material

For all practical purposes each vial contains the same quantity of the substances listed above. **Reconstitute the total contents of one vial of Reagent with 0.5 mL of distilled water.** Allow to stand for a minimum of 5 minutes before use to allow for complete solution of freeze-dried material. Antigen Reagent Lot 2017/118B should be used according to the method described by Wood, JM, Schild, GC, Newman, RW, and Seagroatt, VA, Journal of Biological Standardisation, 1977, 5, 237-247, with the following modification:

It is recommended that Antigen Reagent Lot 2017/118B and test A/Singapore/GP1908/2015 (IVR-180A) virus antigens should be treated with Zwittergent 3-14 detergent (Calbiochem-Behring, La Jolla, CA, USA) before single-radial-immunodiffusion assay. Suitable incubation conditions are as follows:

450 microlitres of antigen are added to 50 microlitres of 10% (w/v) Zwittergent detergent and incubated in covered containers for 30 minutes at room temperature (20-25°C).

Dilutions of detergent treated antigens are then added to wells in single-radial-immunodiffusion plates and incubated at 20-25°C.

Antigen Reagent Lot 2017/118B should be used to assay A/Singapore/GP1908/2015 (IVR-180A) antigens using a TGA antiserum reagent.

No attempt should be made to weigh out any portion of the freeze-dried material. Unopened vials should be store at below -60°C but storage of reconstituted reagent is not recommended. To remove the reconstituted material from the vial, it is necessary to use some form of transfer pipette rather than a volumetric pipette. The contents of the vials should not be assumed to be sterile.

6. Stability

It is the policy of WHO not to assign an expiry date to their international reference materials. They remain valid with the assigned potency and status until withdrawn or amended.

Reference Materials should be stored on receipt as indicated on the label.

**Storage:** Antigen should be stored at or below -60 °C.

**Note:** This material is shipped on dry ice.

The temperature range for shipping is -80 to -70°C.

In addition, once reconstituted, diluted or aliquoted, users should determine the stability of the material according to their own method of preparation, storage and use.

TGA follows the policy of WHO with respect to its reference materials.
7. Citation

In all publications (or data sheets for immunoassay kits) in which this preparation is used as an assay calibrant, it is important that the title of the preparation, vial code and the name and address of TGA are cited correctly.

8. Product liability

Information emanating from TGA is given after the exercise of all reasonable care and skill in its compilation, preparation and issue, but is provided without liability in its application and use.

This product is intended for use as a standard or reference material in laboratory work in relation to biological research, manufacturing or quality control testing of biological products or in the field of *in vitro* diagnostics. It is the responsibility of the user to ensure that he/she has the necessary technical skills to determine the appropriateness of this product for the proposed application. Results obtained from this product are likely to be dependent on conditions of use and the variability of materials beyond the control of TGA.

TGA accepts no liability whatsoever for any loss or damage arising from the use of this product, whether loss of profits, or indirect or consequential loss or otherwise, including, but not limited to, personal injury other than as caused by the negligence of TGA. In particular, TGA accepts no liability whatsoever for:

(i) results obtained from this product; and/or

(ii) non-delivery of goods or for damages in transit.

In the event of any replacement of goods following loss or damage a customer accepts as a condition of receipt of a replacement product, acceptance of the fact that the replacement is not to be construed as an admission of liability on TGA's behalf.
9. Material Safety Sheet

<table>
<thead>
<tr>
<th>Physical properties (at room temperature)</th>
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</thead>
<tbody>
<tr>
<td>Physical appearance:</td>
</tr>
<tr>
<td>White powder</td>
</tr>
<tr>
<td>Fire hazard:</td>
</tr>
<tr>
<td>None</td>
</tr>
</tbody>
</table>

**Chemical properties**

<table>
<thead>
<tr>
<th>Stable:</th>
<th>Yes</th>
<th>Corrosive:</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hygroscopic:</td>
<td>No</td>
<td>Oxidising:</td>
<td>No</td>
</tr>
<tr>
<td>Flammable:</td>
<td>No</td>
<td>Irritant:</td>
<td>No</td>
</tr>
<tr>
<td>Other (specify):</td>
<td>Contains inactivated human influenza virus</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Handling: *See caution, section 4*

**Toxicological properties**

| Effects of inhalation: | No adverse effects have been reported |
| Effects of ingestion: | No adverse effects have been reported |
| Effects of skin absorption: | No adverse effects have been reported |

**Suggested First Aid**

| Inhalation: | Seek medical advice |
| Ingestion: | Seek medical advice |
| Contact with eyes: | Wash with copious amounts of water. Seek medical advice |
| Contact with skin: | Wash thoroughly with water. |

**Action on Spillage and Method of Disposal**

*Spillage of vial contents should be taken up with absorbent material wetted with a virucidal agent. Rinse area with a virucidal agent followed by water. Absorbent materials used to treat spillage should be treated as biologically hazardous waste.*

**Issue Date:** AUGUST 2017