WHO International Standard
1st International Standard for Infliximab
NIBSC code: 16/170
Instructions for use
(Version 2.01, Dated 15/01/2019)

1. INTENDED USE
The World Health Organization (WHO) Expert Committee on Biological standardization (ECBS) recognised the need for a reference standard to evaluate the performance of in vitro biological assays for infliximab.

The preparation 16/170 was assessed in an international collaborative study (described in section 3), and formally adopted by ECBS as the 1st WHO International Standard (IS) for in vitro biological activities of infliximab. In addition, the standard was also recommended by the ECBS for use in therapeutic drug monitoring.

The standard is intended to support the calibration, characterisation and validation of assays used for assessing infliximab and to support the establishment of in-house standards.

It should be noted that the unitage or mass content of the standard should not be used to define the specific activity of infliximab products for regulatory purposes nor to describe product labelling or dosage requirements. Furthermore, the standard and its unitage is not intended to serve any regulatory role in defining biosimilarity, and should not be inferred as serving this purpose.

2. CAUTION
This preparation is not for administration to humans or animals in the human food chain

The preparation contains material of human origin, and either the final product or the source materials, from which it is derived, have been tested and found negative for HBsAg, anti-HIV and HCV RNA. 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 should include the wearing of protective gloves and avoiding the generation of aerosols. Care should be exercised in opening ampoules or vials, to avoid cuts.

3. UNITAGE
The preparation has been assigned the following arbitrary unitage per ampoule:

500 international units (IU)* of tumor necrosis factor-alpha (TNF-alpha) neutralising activity.
500 IU of TNF-alpha binding activity.

The infliximab standard was also tested in assays for antibody dependent cellular cytotoxicity (ADCC) and complement dependent cytotoxicity (CDC) activity, however, due to limited data from the collaborative study, a unitage has not been assigned for these activities.

The use of a mass content of 50 micrograms of infliximab per ampoule is recommended for therapeutic drug monitoring assays (see section 4).

*These units are independent of the amount of TNF-alpha used in various bioassays. For details regarding neutralising activity in terms of the 3rd IS for TNF-alpha (coded 12/154), see report referenced in section 9.

It should be noted that the neutralising activity may vary according to the assay format. Therefore, a relationship between the unitage of the WHO IS coded 16/170 and the activity assigned to in-house standards in the assay system in routine use should be established.

Users should also note that the biological activity of TNF-alpha is likely to vary between different suppliers and this should be controlled by use of an appropriate standard (e.g. WHO IS).

The infliximab IS was tested in a multi-centre collaborative study involving 26 laboratories in 14 countries. Participants tested the IS using assays established in-house, and reported results for cytotoxicity, apoptosis, reporter gene, ADCC, CDC and binding assays (see reference in section 9, WHO/BS/2017.2323).

4. CONTENTS
Country of origin of biological material: United Kingdom.
Each ampoule contains the residue after freeze-drying of 1.0 ml of a solution containing:

Infliximab, approximately 50 micrograms
25mM tri-sodium citrate dihydrate
150mM sodium chloride
1.0% human serum albumin

The infliximab protein was expressed in Sp2/0 cells.

5. STORAGE
Unopened ampoules should be stored at -20ºC.
For economy of use, it is recommended that the solution be sub divided into aliquots and stored at -40ºC or below. Avoid repeated thawing/freezing.

Please note: because of the inherent stability of lyophilized material, NIBSC may ship these materials at ambient temperature.

6. DIRECTIONS FOR OPENING
DIN ampoules have an 'easy-open' coloured stress point, where the narrow ampoule stem joins the wider ampoule body. Various types of ampoule breaker are available commercially. To open the ampoule, tap the ampoule gently to collect material at the bottom (labelled) end and follow manufactures instructions provided with the ampoule breaker.

7. USE OF MATERIAL
No attempt should be made to weigh out any portion of the freeze-dried material prior to reconstitution

Dissolve the total contents of the ampoule in 1.0ml of sterile distilled water.
This solution will contain infliximab at a concentration of 500 IU/ml. Use carrier protein where extensive dilution is required.

8. STABILITY
Reference materials are held at NIBSC within assured, temperature-controlled storage facilities and they should be stored on receipt as indicated on the label. It is the policy of WHO not to assign an expiry date to their international reference materials. Accelerated degradation studies have indicated that this material is suitably stable, when stored at -20ºC or below, for the assigned values to remain valid until the material is withdrawn or replaced. These studies have also shown that the material is suitable for shipment at ambient temperature without any effect on the assigned values. Once reconstituted, diluted or aliquoted, users should determine the stability of the material according to their own method of preparation, storage and use. Users who have data supporting any deterioration in the characteristics of any reference preparation are encouraged to contact NIBSC.

9. REFERENCES
This standard was produced under WHO guidelines cited in the WHO Technical Reports Series, No. 932, 2006, Annex 2.
10. ACKNOWLEDGEMENTS
We are thankful to Celltrion, Inc. for their generous donation of the infliximab material used to develop this preparation. We are grateful to all participants of the collaborative study for their contribution in evaluating the candidate preparations.

11. FURTHER INFORMATION
Further information can be obtained as follows;
This material: enquiries@nibsc.org
WHO Biological Standards:
http://www.who.int/biologicals/en/
JCTLM Higher order reference materials:
http://www.bipm.org/en/committees/jc/jctlm/
Derivation of International Units:
http://www.nibsc.org/standardisation/international_standards.aspx
Ordering standards from NIBSC:
http://www.nibsc.org/products/ordering.aspx
NIBSC Terms & Conditions:
http://www.nibsc.org/terms_and_conditions.aspx

12. CUSTOMER FEEDBACK
Customers are encouraged to provide feedback on the suitability or use of the material provided or other aspects of our service. Please send any comments to enquiries@nibsc.org

13. CITATION
In all publications, including data sheets, in which this material is referenced, it is important that the preparation’s title, its status, the NIBSC code number, and the name and address of NIBSC are cited and cited correctly.

14. MATERIAL SAFETY SHEET

### Physical and Chemical properties

| Classification in accordance with Directive 2000/54/EC. Regulation (EC) No 1272/2008: Not applicable or not Physical appearance: Freeze dried powder | Corrosive: No |
| Stable: Yes | Oxidising: No |
| Hygroscopic: No | Irritant: No |
| Flammable: No | Handling: See caution, Section 2 |
| Other (specify): | Contains material of human origin |

### Toxicological properties

| Effects of inhalation: | Not established, avoid inhalation |
| Effects of ingestion: | Not established, avoid ingestion |
| Effects of skin absorption: | Not established, avoid contact with skin |

### 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 ampoule contents should be taken up with absorbent material wetted with an appropriate disinfectant. Rinse area with an appropriate disinfectant followed by water. Absorbent materials used to treat spillage should be treated as biological waste.

### 16. INFORMATION FOR CUSTOMS USE ONLY

- **Country of origin for customs purposes:** United Kingdom
- **Net weight:** 4.6g
- **Toxicity Statement:** Toxicity not assessed
- **Veterinary certificate or other statement if applicable:** Attached: No

### 17. CERTIFICATE OF ANALYSIS

NIBSC does not provide a Certificate of Analysis for WHO Biological Reference Materials because they are internationally recognised primary reference materials fully described in the instructions for use. The reference materials are established according to the WHO Recommendations for the preparation, characterization and establishment of international and other biological reference standards http://www.who.int/bloodproducts/publications/TRS932Annex2_International Biological_standardisation.pdf (revised 2004). They are officially endorsed by the WHO Expert Committee on Biological Standardization (ECBS) based on the report of the international collaborative study which established their suitability for the intended use.

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