

Therapeutic Goods Information (System for Australian Recall Actions) Specification 2013

*Therapeutic Goods Act 1989*

I, JOHN SKERRITT, a delegate of the Minister for Health, make this Specification under subsection 61(5D) of the *Therapeutic Goods Act 1989*.

Dated 8 March 2013

(signed by)

**JOHN SKERRITT**

Delegate of the Minister for Health

1 Name of Specification

 This Specification is the *Therapeutic Goods Information (System for Australian Recall Actions) Specification 2013*

2 Commencement

 This Specification commences on the day after it is registered.

3 Definitions

 In this Specification:

***Act*** means the *Therapeutic Goods Act 1989*.

***Recall action*** means action taken by the Responsible Entity in order to resolve a problem with therapeutic goods supplied in Australia that have, or may potentially have, deficiencies relating to safety, quality, efficacy or presentation. Recall actions may include: the permanent removal of therapeutic goods from supply in the market; the taking of corrective action in relation to therapeutic goods, such as repair, modification, adjustment or relabelling which may take place at the user's or the Responsible Entity's premises or any other agreed location; and, in the case of medical devices that have been implanted into patients, the issuing of a hazard alert containing information for health practitioners on how to manage such patients.

***Register***means the Australian Register of Therapeutic Goods.

***Responsible Entity*** means, in the case of therapeutic goods on the Register, the person in relation to whom the goods are included in the Register or, in the case of therapeutic goods not on the Register, the supplier of those therapeutic goods or any another person who undertakes the recall action.

***TGA*** means the Therapeutic Goods Administration, being a division of the Department of Health and Ageing.

4 Therapeutic goods information

 The kinds of therapeutic goods information mentioned in Schedule 1 are specified under subsection 61(5D) of the Act for the purposes of subsection 61(5C) of the Act.

Schedule 1 Specified kinds of therapeutic goods information

(section 4)

The following kinds of therapeutic goods information in relation to recall actions, being information kept by the TGA in its System for Australian Recall Actions database:

Note: The following specified kinds of therapeutic goods information may be released by the Secretary to the public under subsection 61(5C) of the Act.

1. the recall commencement date, being the date on which the recall action commenced;
2. a product description of the therapeutic goods being recalled including the brand name(s) of the goods, any active ingredients in the goods (where relevant) and any other information that is necessary to identify the goods;
3. the product type, e.g., whether it is a medicine or a medical device;
4. the recall action taken, i.e., whether the therapeutic goods have been or are being permanently removed from the market, whether other corrective action has been, or is being, taken or if a hazard alert has been issued;
5. the class of recall, being Class I, Class II or Class III, with Class I representing the greatest risk and Class III the lowest risk, to safety;
6. the level of recall action being taken, i.e., whether it is a recall of therapeutic goods from hospitals, wholesalers, retailers and/or consumers;
7. the ARTG number of the recalled goods (where applicable), being the identifier number for the therapeutic goods that are registered, listed or included in the Register;
8. the TGA recall reference, being the unique identifier number the TGA gives to the recall action;
9. the Responsible Entity;
10. recall instructions, being instructions given by the TGA or the Responsible Entity to health practitioners and consumers of the therapeutic goods being recalled; and
11. contact information, being information about the person that the customer or consumer should contact for additional information about the therapeutic goods the subject of the recall action (e.g., a representative of the Responsible Entity).

**Note**

1. All legislative instruments and compilations are registered on the Federal Register of Legislative Instruments kept under the *Legislative Instruments Act 2003.* See http://www.frli.gov.au