



Public Summary

Summary for ARTG Entry: 134893 Allegro Concepts Pty Ltd - Lift, patient transfer, sling/harness/strap

ARTG entry for Medical Device Included Class 1

Sponsor Allegro Concepts Pty Ltd

Postal Address 30 James Street, LIDCOMBE, NSW, 2141

Australia

ARTG Start Date 12/02/2007

Conditions

The automatic conditions applicable to the inclusion of all kinds of medical devices in the Register are as specified in section 41FN of the Therapeutic Goods Act 1989.

The standard conditions that are imposed under section 41FO of the Therapeutic Goods Act 1989 when kinds of medical devices are included in the Register are as set out in the following paragraphs.

For a medical device included in the Register under Chapter 4 and imported into Australia, the Sponsor must ensure that information about the Sponsor is provided in such a way as to allow the sponsor to be identified.

Each sponsor shall retain records of the distribution of all of the sponsor's medical devices included in the Register under Chapter 4. In the case of records relating to a Class AIMD medical device, Class III medical device, or Class IIb medical device that is an implantable medical device, the distribution records shall be retained for a minimum period of 10 years. In the case of records relating to any other device, the distribution records shall be retained for a minimum period of 5 years.

The sponsor of a medical device included in the Register under Chapter 4 shall keep an up to date log of information of the kind specified in Regulation 5.8.

The sponsor shall provide to the Director, Office of Devices, Blood and Tissues, Therapeutic Goods Administration, three consecutive reports which include information of a kind specified in Regulation 5.8 that arises during the reporting period, concerning Class III, Class AIMD or implantable Class IIb medical devices. The reporting period for the first report commences on the date of inclusion and is to be at least a period of six months but not more than 18 months ending on 1 October. For subsequent reports the reporting period is 12 months ending on 1 October.

Where a medical device included in the Register, contains a substance which is included in the Fourth Schedule to the Customs (Prohibited Imports) Regulations or the Eighth Schedule to the Customs (Prohibited Exports) Regulations the Sponsor shall, at the time of importation or exportation of the medical device, be in possession of a licence and a permission for importation or exportation of each consignment of the goods as required by those regulations.

A sponsor shall ensure that a medical device within their control is stored and transported in accordance with the instructions and information provided by the manufacturer.

Manufacturers

NameAddressStepsAllegro Concepts Pty Ltd - 4858430 James StreetManufacturing steps not recorded.

LIDCOMBE, NSW, 2141

Products

1. Lift, patient transfer, sling/harness/strap

Product Type Single Device Product Effective date 12/02/2007

GMDN 40535 Lift, patient transfer, sling/harness/strap

Functional description Not included on record

Intended purpose A sling is an item of moving and handling equipment. It is used with a mechanical hoist to facilitate the

lifting and transfering of clients.

Variant information