# Manufacturer's Declaration of Conformity for Class I non-sterile, non-measuring or Class 1 in vitro diagnostic (IVD) medical devices

This Declaration of Conformity (DoC) is required under clause 6.6 (for single devices or kinds of devices) of Schedule 3 of the <u>Therapeutic Goods (Medical Devices) Regulations 2002</u> (the Regulations).

An Australian DoC must be completed by the manufacturer of a Class I non-sterile, non-measuring device or a Class 1 in vitro diagnostic (IVD) device.

For more information on how to complete this DoC refer to <u>Guidance for Declaration of Conformity for Class I non-sterile non-measuring and Class 1 in vitro diagnostic (IVD) medical devices.</u>

This document can be:

- used for single or multiple devices.
- filled out by hand and then scanned and submitted, or filled out electronically.

### Manufacturer's details

Manufacturer's name	Qlicksmart Pty Ltd
Manufacturer's business address	Level 1, Boundary Street, West End, Brisbane, QLD 4101, Australia

# **Classification type**

Specify if your device is:

	Class	I non-sterile	non-measuring	device
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Class 1 in vitro diagnostic (IVD) medical device

### **GMDN** code and term

Select the most appropriate Global Medical Device Nomenclature (GMDN) code for this product. GMDN codes and terms are a system of internationally agreed generic descriptors that are used to identify all medical device products. Class 1 IVDs require the use of a level 1 collective term (CT). Please refer to the following link for guidance regarding an appropriate CT for the kind of device: The use of GMDN codes for IVD medical devices in Australia.

GMDN codes are generated by the GMDN Agency.

The GMDN code tables are available on TGA Business Services (TBS).

GMDN codes	46240, Syringe/ampoule clip
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### Standards applied to the device(s)

List any standards used in the manufacturing of the device, including:

- International Standards (ISO)
- Australian Standards (AS)
- Conformity Assessment Standard Orders (CASO)
- Medical Device Standard Orders (MDSO)

Standards	EN ISO 15223-1:2021 Medical devices – Symbols to be used with medical device labels, labeling and information to be supplied Part 1: General requirements
	EN ISO 14971:2019 Medical devices – Application of risk management to medical devices
	EN ISO 20417:2021 Medical Devices - Information to be supplied by the manufacturer

# Name of medical device(s) / IVD(s)

Identify the specific device/s this form relates to for example, the name of the device or model numbers of the device and any variants that are being manufactured for supply in Australia.

Name of medical device(s)/IVD(s)	Qlicksmart CheckCLIP
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This declaration is being made under clause 6.6 of Schedule 3 to the Regulations for a Class I non-sterile, non-measuring device or Class 1 IVD medical device.

## By signing this form you are agreeing that:

- You have reviewed the Declaration of Conformity procedures under Part 6 of Schedule 3 of the Regulations and the device complies with the applicable provisions of those procedures.
- The device complies with the relevant essential principles set out in Schedule 1 to the Regulations.
- The device complies with the applicable provisions of the classification rules set out in Schedule 2 to the Regulations.
- You will update the technical documentation when any changes are made in relation to the device.



### Important note

Providing information that is false or misleading to a Commonwealth entity or in connection with a Commonwealth law is a serious offence subject to criminal penalties under the *Criminal Code Act 1995*.

Name	Michael Sinnott
Title	Managing Director
Signature	Date 01/11/2022