



Declaration of Conformity

According to Directive 98/79/EC on In-Vitro Diagnostic Medical Devices, Annex III

Manufacturer: Glenbio Ltd.
10 Kilbegs Road
Antrim
Co. Antrim
BT41 4NN
United Kingdom

Product: EC40, EZ-CYTO Thin Layer Processor

Classification: General category in-vitro diagnostic medical device.
[Other Device (all devices except Annex II and self-testing devices)].

We hereby declare that the requirements imposed by the IVD Directives 98/79/EC and the Statutory Instruments 2002 No. 618 (Consolidated legislation) have been fulfilled.

Signed

Date

Amy McDonnell

17 March 2017

Amy Mc Donnell
The QA/RA Manager



Declaration of Conformity

According to Directive 98/79/EC on In-Vitro Diagnostic Medical Devices, Annex III

Manufacturer: Glenbio Ltd.
10 Kilbegs Road
Antrim
Co. Antrim
BT41 4NN
United Kingdom

Product: EZ-CYTO KIT 200 TESTS

Classification: General category in-vitro diagnostic medical device.
[Other Device (all devices except Annex II and self-testing devices)].

We hereby declare that the requirements imposed by the IVD Directives 98/79/EC and the Statutory Instruments 2002 No. 618 (Consolidated legislation) have been fulfilled.

Signed

Date



17 March 2017

Amy Mc Donnell
The QA/RA Manager



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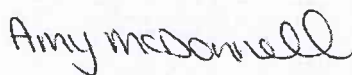
Product: EZ-CYTO KIT 400 TESTS

Classification: General category in-vitro diagnostic medical device.
[Other Device (all devices except Annex II and self-testing devices)].

We hereby declare that the requirements imposed by the IVD Directives 98/79/EC and the Statutory Instruments 2002 No. 618 (Consolidated legislation) have been fulfilled.

Signed

Date



17 March 2017

Amy Mc Donnell
The QA/RA Manager