

MORE DIAGNOSTICS, INC.

Declaration of Conformity

We hereby declare that the in vitro diagnostic devices and the accessories described below conform to all of the applicable essential requirements contained in the Annex I Essential Requirements of In Vitro Diagnostic Device Directive 98/79/EC as published in the European Official Journal on 1999-12-07.

This conformance is supported by the technical documentation which is maintained by MORE DIAGNOSTICS, Inc.

Product: Cyclosporine C2 Control

Catalog Number (REF): 202 - Tri-level package

Manufacturer Address: More Diagnostics, Inc.
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Los Osos, CA 93402

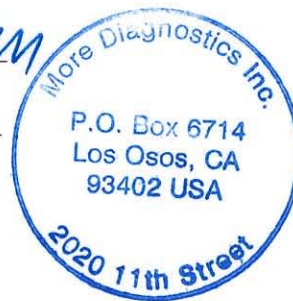
EC Authorized Representative: Siemens Healthcare Diagnostics Limited
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Date: 2018-3-26

Authorization:

Signature Harry R. Holden

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RA/QA Management Representative



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