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Product Service

Verification Report

according to Directive 98/79/EC Annex IV.6

Choose certainty.
Add value.**No.** ROC-05390095 17 12 040365**Manufacturer:** ROCHE Diagnostics GmbH
Sandhofer Strasse 116
D-68305 MannheimMat-Nr.: 05390095190
HIV combi PT Elecsys cobra
Batch: 28316801
2~8°C 
Lot: 040000206248 17.11.2017
Sample No./Type: 200137985/Y2**Product:** HIV Combi PT
Elecsys and cobas e analyzers**Batch:** 28316801**Expiry Date:** 06-2018

The above mentioned batch meets the batch release criteria established during design examination and may be placed on the market. The design examination certificate issued for this product is V7 16 10 10283 366.

Date, 2017-12-04
i.A. Dr. Laura Scrivano
In-vitro-Diagnostika

TÜV SÜD Product Service GmbH is Notified Body according to Council Directive 98/79/EC concerning In Vitro Diagnostic Medical Devices with Identification No. 0123.

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