

## EU DECLARATION OF CONFORMITY for REAGENTS FOR DIATABS

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ROSCO DIAGNOSTICA A/S hereby declares that the following products:

AMINOPEPTIDASE REAGENT

NINHYDRIN 3.5 % SOLUTION and N,N-DIMETHYL- $\alpha$ - NAPHTHYL

KOVAC's REAGENT and SULFANILIC ACID 0.8 % SOLUTION,

comply with the In Vitro Diagnostic Medical Devices Directive 98/79/EC.


According to the directive the above reagents are classified in the group of 'Other Devices', i.e. devices not listed in List A or B in Annex II and devices not intended for Performance Evaluation. Conformity route: Annex III.

The Declaration covers the above reagents (DVL0005) distributed from ROSCO DIAGNOSTICA A/S, which have been supplied with a CE-mark for compliance.

Date of Validity:

04.02.2009

Authorization:

  
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QA Manager

ROSCO DIAGNOSTICA's List of Reagents = DVL0005 is available on request, and shall continuously be updated.