

## **EC DECLARATION OF CONFORMITY**

We,

**Thermo Fisher Scientific Oy, Clinical Diagnostics Finland**

Hereby declare that the Selective Clinical Chemistry Analyzers with

Trade name:	<b>Konelab PRIME 30</b>	<b>Konelab PRIME 30 ISE</b>
Type:	<b>986</b>	<b>987</b>

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

This Declaration is valid for all devices which are placed on the market by ourselves on or after October 20, 2008 and which bear the CE marking.

Vantaa, November 04, 2008

Thermo Fisher Scientific Oy



Silja Halme  
Quality Manager  
Quality, Regulatory and Compliance  
Clinical Diagnostics Finland