

## EU DECLARATION OF CONFORMITY

## Manufacturer:

DiaSys Diagnostic Systems GmbH Alte Strasse 9 65558 Holzheim Germany

## SRN: DE-MF-000025417

Product name	Cat. No.	Basic UDI-DI	Intended use	Risk class
respons®940	965 400	4051839965400ZM	Random access clinical chemistry analyzer for in vitro diagnostic use. For professional use only.	A
respons®940 complete	965 900	4051839965400ZM	Random access clinical chemistry analyzer for in vitro diagnostic use. For professional use only.	A
respons®940 (ISE)	965 500	4051839965500ZS	Random access clinical chemistry analyzer with integrated ion selective electrodes unit (ISE) for in vitro diagnostic use. For professional use only.	A
respons®940 complete (ISE)	965 910	4051839965500ZS	Random access clinical chemistry analyzer with integrated ion selective electrodes unit (ISE) for in vitro diagnostic use. For professional use only.	A
Cleaner A	186109910923	04051839186109C	Wash solution for clinical chemistry analyzer	А
Cleaner B	186509910923	04051839186509Q	Wash solution for clinical chemistry analyzer	А
Cleaner respons 920/940	188309910885	04051839188309U	Wash solution for clinical chemistry analyzer	А

We, as the manufacturer of the devices take sole responsibility for and hereby declare that the mentioned products listed above meet the provisions of the Regulation (EU) 2017/746 on invitro diagnostic medical devices.



Conformity route:

Annex IX Quality Management System for class A devices and technical documentation according Annex II and Annex III.

Additional applicable directives: Directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS).

DiaSys Diagnostic Systems GmbH Holzheim, 2023-01-23 Dr. Jan Gorka Managing Director + CEO

CE