


DECLARATION OF CONFORMITY



Product name / Trade name		REF IS-4530NG	BASIC UDI-DI: 5060169696457YN				
IDS ACTH II Control Set		GMDN: 41796	EMDN: W0102152008				
Intended Purpose	For In Vitro Diagnostic Use For Laboratory Professional Use The IDS ACTH II Control Set is for in vitro diagnostic use, for the quality control of the IDS ACTH II assay on the IDS system.						
 Immunodiagnostic Systems Limited, 10 Didcot Way, Boldon Business Park, Boldon, Tyne and Wear, NE35 9PD, UK		<table border="1"><tr><td>EC</td><td>REP</td></tr><tr><td colspan="2">Immunodiagnostic Systems SA 101 Rue Ernest Solvay B-4000 Liege Belgium</td></tr></table> SINGLE REGISTRATION NUMBER: GB-MF-000015851		EC	REP	Immunodiagnostic Systems SA 101 Rue Ernest Solvay B-4000 Liege Belgium	
EC	REP						
Immunodiagnostic Systems SA 101 Rue Ernest Solvay B-4000 Liege Belgium							
		SINGLE REGISTRATION NUMBER: BE-AR-000015342					

RISK CLASS: ☐ A ☒ B ☐ C ☐ D

CLASSIFICATION RULE (ANNEX VIII) : Rule 6

CONFORMITY ROUTE: ☒ ANNEX IX Full Quality System
(Class B, C & D)

☐ ANNEX I & II+III
(non-sterile Class A)

Name of the Notified Body / Identification: TÜV Rheinland LGA Products GmbH.
0197

CE Marking Date: 30th October 2023

EU CERTIFICATE No.: HX 2333083-1

COMMON SPECIFICATIONS: Not Applicable

We, as the manufacturer of the device(s) take sole responsibility for and hereby declare that the above-mentioned product(s) meet(s) the provisions of the following Regulation(s)/Directives

- Regulation EU 2017/746 on *In vitro* Diagnostic Medical Devices

Date: 15th April 2024

Place: UK

Signed on behalf of Immunodiagnostic Systems Limited


M Henderson
RA Manager & PRRC

Date: 15th April 2024

Place: UK

Signed on behalf of Immunodiagnostic Systems Limited


D. Mullington
Group QA/RA Director & PRRC