

DECLARATION OF CONFORMITY

Manufacturer:

Invivoscribe, Inc.

10222 Barnes Canyon Rd. Bldg 1 San Diego, California 92121 United States of America

Authorized Representative:

Invivoscribe Technologies, SARL

c/o Ficorec Domiciliation Services

132. Boulevard Michelet Hall Nord - 5ème étage

13008 Marseille

FRANCE

UK Responsible Person:

Gillian Pawlowsky Ltd.

272 Bath Street Glasgow G2 4JR Scotland UK

Family Name:

LymphoTrack® Dx

Device Trade Name:

IGH FR2 Assay Kit - MiSeq

Catalog #	Device	Quantity	UDI-DI	GTIN	EMDN Code
9-121-0089	LymphoTrack® Dx IGH FR2	40	08100227391210089R6	850052003838	W010699
	Assay Kit A – MiSeq®	Reactions			
9-121-0099	LymphoTrack® Dx IGH FR2	120	08100227391210099R9	850052003845	W010699
	Assay Panel – MiSeq®	Reactions			

I, the undersigned, hereby declare that the in-vitro diagnostic medical devices specified above conform to the European Directive 98/79/EC, In vitro Diagnostic Medical Device Directive, Annex III.

I, the undersigned, hereby declare that the in-vitro diagnostic medical devices specified above conform to the Part IV of the UK Medical Device Regulations 2002, Annex III (as modified by Part III of Schedule 2A to the UK MDR 2002).

*Date of Validity: 10 Oct Z0Z3

Jason Gerhold

Global Director of Quality, Regulatory and Clinical Affairs

Invivoscribe, Inc.

10222 Barnes Canyon Rd. Bldg 1

San Diego, California 92121

USA

^{*}Originally signed on 05/17/2022, no significant changes have occurred to the product since that date