



DECLARATION OF CONFORMITY

Manufacturer: **Invivoscribe Inc.**
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United States of America

Authorized Representative: **Invivoscribe Technologies, SARL**
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
Family Name: LymphoTrack® Dx

Device Trade Name: **TRG Assay – S5/PGM**

Catalog #	Device	Quantity
9-227-0007	LymphoTrack® Dx TRG Assay – S5/PGM	60 Reactions

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

Date of Validity: 13 Feb 2019

By: 

Jason Gerhold
Global Director of Regulatory Affairs & Quality Assurance
Invivoscribe Inc.
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