

CERTIFICATE OF GMP COMPLIANCE

We certify herewith

that the company Suisse Technology Partners AG, Querstrasse 5, 8212 Neuhausen am Rheinfall, Authorisation No. 512265-102645393 with its site Suisse Technology Partners AG, Querstrasse 5, 8212 Neuhausen am Rheinfall, Switzerland, Site No. 1003073 has been duly authorised to perform the manufacturing activities according to the table below;

that the company is keeping the required level for Good Manufacturing Practices for Medicinal Products (GMP) according to the Swiss regulations in force. These regulations are in accordance with the requirements for good practices in the manufacture and quality control of the Pharmaceutical Inspection Convention/Co-operation Scheme (PIC/S) as well as with The Good Manufacturing Practice requirements referred to in the Agreement of Mutual Recognition between the European Union/Canada and Switzerland;

No.	Operation	Scope
1	MANUFACTURE OF MEDICINAL PRODUCTS (WITHOUT LABILE BLOOD PRODUCTS)	
1.6	Quality control testing	
1.6.3	Chemical/Physical	H/V
3	MANUFACTURE OF ACTIVE SUBSTANCES	
3.6	Quality control testing of medicinal products	
3.6.1	Physical / Chemical testing	-

that the manufacturing plant of the company is subject to official periodic inspections; the last regular inspection was conducted on **20.10.2020** (dd.mm.yyyy).

* Scope of authorisation:

H/V Human and veterinary medicinal products, without investigational products

V Veterinary medicinal products only, without investigational products

I Human investigational medicinal products

Not specified

Berne, 07.01.2021 (dd.mm.yyyy) No. GMP-CH-1001752



Swissmedic, Swiss Agency for Therapeutic Products

M. Bar

Marianne Baumann

Schweizerisches Heilmittelinstitut Institut suisse des produits thérapeutiques Istituto svizzero per gli agenti terapeutici Swiss Agency for Therapeutic Products

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