

CERTIFICATE OF GMP COMPLIANCE

We certify herewith

that the company **Bühlmann Laboratories AG, Baselstrasse 55, 4124 Schönenbuch, Switzerland**, has been duly authorized to manufacture and distribute medicinal products (in-vitro testsystems/bio assays);

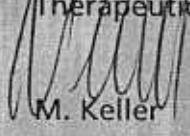
that the company is keeping the required level for good practices in the manufacture of medicinal products according to the Swiss regulations in force. These regulations are in accordance with the requirements for good practices in the manufacture and quality control of the Pharmaceutical Inspection Convention /Co-operation Scheme (PIC/S) and the Directives of the European Commission;

that the manufacturing plant of the company is subject to official periodic inspections; the last inspection was conducted on 14 February 2001;

that the requirements regarding manufacture and quality control for medicinal products for export are identical to those applicable to products sold in Switzerland.

Bern, February 8, 2002
No. HK02-083

Swissmedic, Swiss Agency for
Therapeutic Products


M. Keller



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