



ORSZÁGOS GYÓGYSZERÉSZETI INTÉZET
NATIONAL INSTITUTE OF PHARMACY

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HUNGARY

Budapest, 05.08.2003
No.: 1127 -100/38/2003
Our ref
Annex
Subject: Italy

CERTIFICATE

It is hereby certified that the manufacturing plant of

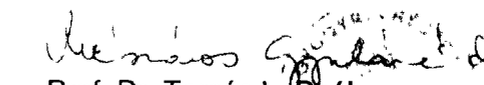
Biropharma First Hungarian Biotechnological Ltd. 6413 Kunfehértó, IV. körzet 6.

is authorized for the manufacture and sale of pharmaceutical products and preparations in Hungary.

In accordance with 25th Act of 1998 on medicinal products for human use, Biropharma Ltd. is subject to regular inspections made by the National Institute of Pharmacy in respect of Good Manufacturing Practices and Quality Control according to the recommendations of the World Health Organization and requirements of the Pharmaceutical Inspection Convention (PIC), and Pharmaceutical Cooperation Scheme (PIC/S).

(Mandatory GMP compliance for medicine manufacturers was originally introduced by the 31st Law-Decree of 1976.)

The certificate is valid for two years.


for Prof. Dr. Tamás L. Paál
Director General

Certificate Q17677



The management system of

Biropharma Kft.

6413 Kunfehértó
IV körzel 6



has been assessed and certified as meeting the requirements of

ISO 9001:2000

For the following activities

Development, manufacture and contract manufacture of dietary supplements, nutraceuticals, phytopharmaceuticals and drug intermediates.

Further clarifications regarding the scope of this certificate and the applicability of ISO 9001 2000 requirements may be obtained by consulting the organization

This certificate is valid from 9 April 2003 until 15 March 2006.
Issue 2. Certified since March 2000.

Authorised by



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