



**European Directorate for the Quality of Medicines
Division Certification of Substances**

**Certificate of suitability
No. R1-CEP 2001-030-Rev 00**

Name of the substance:

FOETAL BOVINE SERUM

Product codes S 0313/ S 0315/ S 0210/ S 0213/ S 0215

Name of holder:

BIOCHROM AG

Leonorenstrasse 2-6

D - 12247 Berlin

Site of production:

BIOCHROM AG

Leonorenstrasse 2-6

D - 12247 Berlin

**THIS CERTIFICATE SUPERSEDES THE PREVIOUS CERTIFICATE
R0-CEP 2001-030-REV 02**

After examination of the information provided on the origin of raw material(s) and type of tissue(s) used and on the manufacturing process for this substance on the site of production mentioned above, D - 12247 Berlin, we certify that the substance **FOETAL BOVINE SERUM** meets the criteria described in the current version of the monograph Products with risk of transmitting agents of animal spongiform encephalopathies no. 1483 of the European Pharmacopoeia, current edition including supplements.

- country(ies) of origin of source materials: United States of America and Argentina
- nature of animal tissues used in manufacture: Foetal bovine blood

The submitted dossier must be updated after any significant change that may alter the quality, safety or efficacy of the substance, or that may alter the risk of transmitting animal spongiform encephalopathy agents.

Manufacture of the substance shall take place in accordance with a suitable quality assurance system such as GMP, and in accordance with the dossier submitted.

Failure to comply with these provisions will render this certificate void.

The certificate is valid provided that there has been no deterioration in the TSE status of the country(ies) of origin of the source material.

- 30 This certificate is renewed from **7 December 2006** according to the provisions of Resolution
31 AP-CSP (93) 5 as amended, and of Directive 2001/83/EC and Directive 2001/82/EC and any
32 subsequent amendment, and the related guidelines.
33 This certificate has 33 lines only.



Dr. A. ARTIGES
Director of the Quality of Medicines

Strasbourg, 13 November 2006

DECLARATION OF ACCESS *(to be filled in by the certificate holder under their own responsibility)*

BIOCHROM AG, as holder of the certificate of suitability

R1-CEP 2001-030-Rev 00 for FOETAL BOVINE SERUM

hereby authorises
(name of the pharmaceutical company)

to use the above-mentioned certificate of suitability in support of their application(s) for the following Marketing
Authorisation(s): *(name of product(s) and marketing number(s), if known)*

The holder also certifies that no significant changes to the operations as described in the CEP dossier have been
made since the granting of this version of the certificate.

Date and Signature *(of the CEP holder)*: