

Certification of Substances Department

Certificate of suitability
No. R0-CEP 2016-320-Rev 00

1 *Name of the substance:*

2 **CELECOXIB**

3 *Name of holder:*

4 **SCINOPHARM TAIWAN, LTD.**

5 No. 1, Nan-Ke 8th Road

6 Taiwan-74144 Shan-Hua, Tainan

7 *Site(s) of production:*

8 **SEE ANNEX 1**

Notice

NOT FOR REGISTRATION PURPOSES

**For filing purposes please contact ScinoPharm Taiwan
to obtain a complete "controlled copy" of this CEP.**

**ScinoPharm Taiwan- Regulatory Technical Services
(SPT.RTS@scinopharm.com.tw)**

subsequent
annex, we
on of the
including
analytical

(Annex 2)

18 – Test for the following impurity by liquid chromatography (Annex 3)

19 4-Sulfonamidophenylhydrazine hydrochloride

20 (4-SAPH)

not more than 3.75 ppm

21 – Test for residual solvents by gas chromatography (Annex 4)

22 Isopropanol

not more than 5000 ppm

23 – Test for residual solvents by liquid chromatography (Annex 5)

24 Trifluoroacetic acid

not more than 3000 ppm

25 In the last steps of the synthesis water is used as solvent.

26 No elemental impurity classified in ICH Q3D is intentionally introduced in the manufacture of
27 the substance.

28 The re-test period of the substance is 12 months if stored in a double polyethylene bag, in an
29 aluminium foil bag placed in a polyethylene drum.

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F-67081 Strasbourg (France)

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Internet: <http://www.edqm.eu>

30 The holder of the certificate has declared the absence of use of material of human or animal
31 origin in the manufacture of the substance.

32 The submitted dossier must be updated after any significant change that may alter the quality,
33 safety or efficacy of the substance.

34 Manufacture of the substance shall take place in accordance with the Good Manufacturing Practice
35 and in accordance with the dossier submitted.

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

37 This certificate is granted within the framework of the procedure established by the European
38 Pharmacopoeia Commission [Resolution AP-CSP (07) 1] for a period of five years starting from
39 **18 December 2017**. Moreover, it is granted according to the provisions of Directive 2001/83/EC
40 and Directive 2001/82/EC and any subsequent amendment, and the related guidelines.

41 This certificate has five annexes, the first of 1 page, the second, the third and the fourth of 3
42 pages each and the fifth of 2 pages.

43 This certificate has:
44 lines.



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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):