





National Authority of Medicines and Health Products, I.P.

CERTIFICATE NUMBER: FT069/MH/001/2018

CERTIFICATE OF GMP COMPLIANCE OF A MANUFACTURER

Part 1

Issued following an inspection in accordance with:

Art. 111(5) of Directive 2001/83/EC as amended

The competent authority of Portugal confirms the following:

The manufacturer: Taha Pharma Laboratoire Pharmaceutique

Site address: Zone Industrielle de Medjez, Lote 14,, Meddjez El Bab,, Beja, 9070, Tunisia

Has been inspected in connection with marketing authorisation(s) listing manufacturers located outside of the European Economic Area in accordance with Art. 111(4) of Directive 2001/83/EC transposed in the following national legislation:

Art.176.º n.º 4 of Decree-Law n.º 176/2006, 30 of August

From the knowledge gained during inspection of this manufacturer, the latest of which was conducted on *2018-02-09*, it is considered that it complies with:

The principles and guidelines of Good Manufacturing Practice laid down in Directive 2003/94/EC

This certificate reflects the status of the manufacturing site at the time of the inspection noted above and should not be relied upon to reflect the compliance status if more than three years have elapsed since the date of that inspection. However, this period of validity may be reduced or extended using regulatory risk management principles by an entry in the Restrictions or Clarifying remarks field. This certificate is valid only when presented with all pages and both Parts 1 and 2. The authenticity of this certificate may be verified in EudraGMDP. If it does not appear, please contact the issuing authority.

Pagernando Direcção Inspecção e Licenciamentos

EudraGMDP, Ref key: 50022

Issuance Date: 2018-09-11

Signatory: Ms. M. F. R. H. Matos

¹ The certificate referred to in paragraph 111(5) of Directive 2001 83/EC and 80(5) of Directive 2001/82/EC, shall also be required for imports coming from third countries into a Member State.

Guidance on the interpretation of this template can be found in the Help menu of EudraGMDP database.

³ These requirements fulfil the GMP recommendations of WHO.







Part 2

Human Medicinal Products

1.2	Non-sterile products
	1.2.1 Non-sterile products (processing operations for the following dosage forms)
	1.2.1.1 Capsules, hard shell
	1.2.1.13 Tablets
1.5	Packaging
	1.5.1 Primary Packing
	1.5.1.1 Capsules, hard shell
	1.5.1.13 Tablets
	1.5.2 Secondary packing
1.6	Quality control testing
	1.6.2 Microbiological: non-sterility
	1.6.3 Chemical/Physical

2018-09-11

Name and signature of the authorised person of the Competent Authority of Portugal

Ms. Maria Fernanda Ralha Henriques Maternanda Ralha
National Authority of Maria National Authority of Medicines and Health Products cenciamentos
I.P. Inspecção

Tel: +351 21 7987278 Fax: +351 21 7987257



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Signatory: Ms. M. F. R. H. Matos