MANUFACTURER'S AUTHORISATION

1. Authorisation Number 0752

2. Name of authorisation holder Uždaroji akcinė bendrovė "NORAMEDA"

3. Address(es) of manufacturing site(s) Uždaroji akcinė bendrovė "NORAMEDA", Gynėjų g. 16, Vilniaus

m. sav., Vilniaus m., LT-01109, Lithuania

4. Legally registered address of authorisation

holder

Meistrų g. 8A, Vilniaus m. sav., Vilniaus m., LT-02189, Lithuania

5. Scope of authorisation and dosage forms ² ANNEX 1 and/ or ANNEX 2

6. Legal Basis of authorisation Art. 40 of Directive 2001/83/EC

7. Name of responsible officer of the competent authority of the member state granting the

manufacturing authorisation

confidential

8. Signature

9. Date 2006-05-25

10. Annexes attached Annex 1 and/or Annex 2

Optional Annexes as required:

Annex 3 (Addresses of Contract Manufacturing Site(s))

Annex 4 (Addresses of Contract laboratories)

Annex 5 (Name of Qualified Person)

Annex 6 (Name of responsible persons)

Annex 7 (Date of inspection on which authorisation granted, scope of last

inspection)

Annex 8 (Manufactured/imported products authorised)³

Online EudraGMDP, Ref key: 4770 MANUFACTURER'S AUTHORISATION: 0752

¹ The authorisation referred to in paragraph 40(1) of Directive 2001/83/EC and 44(1) of Directive 2001/82/EC, as amended, shall also be required for imports coming from third countries into a Member State.

² Guidance on the interpretation of this template can be found in the Help menu of EudraGMDP database.

³ The Competent Authority is responsible for appropriate linking of the authorisation with the manufacturer's application (Art. 42(3) of Directive 2001/83/EC and Art. 46(3) of Directive 2001/82/EC as amended).

SCOPE OF AUTHORISATION

ANNEX 1

Name and address of the site: Uždaroji akcinė bendrovė "NORAMEDA", Gynėjų g. 16,

Vilniaus m. sav., Vilniaus m., LT-01109, Lithuania

Human Medicinal Products

Authorised Operations

MANUFACTURING OPERATIONS (according to part 1)
IMPORTATION OF MEDICINAL PRODUCTS (according to part 2)

Part 1 - MANUFACTURING OPERATIONS	
1.1	Sterile products
	1.1.3 Batch certification
1.2	Non-sterile products
	1.2.2 Batch certification
Part 2	2 - IMPORTATION OF MEDICINAL PRODUCTS
2.2	Batch certification of imported medicinal products
	2.2.1 Sterile products
	2.2.1.2 Terminally sterilised
	2.2.2 Non-sterile products