

From :  
The Drugs Controller General (India)  
Directorate General of Health Services

FDA Bhawan, New Delhi

Dated 28 APR 2011

To,

M/S. B.M CHEMIE  
BLDG.NO.B,GALA NO.110 ,1ST FLOOR  
,SHREE SAMBHAV COMPLEX,REHNA  
VILLAGE,BHIWANDI, DISTRICT-THANE  
MAHARASHTRA

Sub: -Registration of M/S. HENAN LIHUA PHARMACEUTICAL CO. LTD. MIDDLE OF HUANGHE STREET, ANYANG HI-TECH INDUSTRY DEVELOPMENT ZONE, HENAN CHINA under the Provisions of Drug & Cosmetics Rules for the purpose of import of drugs in India

Dear Sir,

Please refer to your application No. NIL dated 04/02/2011: received by this Office vide diary No : 7224 dated : 15/02/2011 on the above subject. Registration Certificate in Form 41 under the Rules is issued for the manufacturing site along with the name (s) of drug (s) imported under the said Certificate subject to the condition:

1. The drugs shall conform to the standards / specifications mentioned in the Second Schedule of the Act, or such other standards / specifications forwarded by you and approved by this Directorate.
2. The drugs shall have the composition, strengths, standards and packing as approved by this Directorate.
3. Dispute, if any, in respect of the payment of fees and submission of TR6 challan, shall have to be settled between the bank and the applicant.
4. The drugs will be required to be withdrawn from sale from the market in case any undesirable reactions due to its medication are brought to light at any stage. This Directorate should be informed of adverse reports on the drug, if any.
5. The drugs shall be marketed for indications for which they have been approved.
6. If any claim is made for these drugs in future other than those approved, prior permissions from this Directorate will be necessary before such claim is made.
7. The drugs shall be supplied and marketed as per the provisions of the existing Drugs & Cosmetics Act and Rules thereunder.

Conditions of the Registration Certificate

1. The Registration Certificate shall be displayed at a prominent place by the authorised agent.
2. No drug shall be registered unless it has a free sale approval in the country of origin, and/or in other major countries.
3. The manufacturer or his authorised agent in India shall comply with the conditions of the import licence issued under the Drugs and Cosmetics Rules, 1945.
4. The manufacturer or his authorised agent in India shall inform the licensing authority forthwith in the event of any administrative action taken due to adverse reaction, viz. market withdrawal, regulatory restrictions, or cancellation of authorisation, and/or not of standard quality report of any drug pertaining to this Registration Certificate declared by the Regulatory Authority of the country of origin or by any Regulatory Authority of any other country, where the drug is marketed/sold or distributed. The despatch and marketing of the drug in such cases shall be stopped immediately, and the licensing authority shall be informed immediately. Further action in respect of such stopped marketing of drug shall be followed as per the direction of the licensing authority. In such cases, action equivalent to that taken with reference to the concerned drug in the country of origin or in the country of marketing shall be followed in India also, in consultation with the licensing authority. The licensing authority may, however, direct any further modification to this course of action, including the withdrawal of the drug from Indian market within 48 hours time period.
5. The manufacturer or his authorised agent in India shall inform the licensing authority within 30 days in writing in the event of any change in manufacturing process, or in packaging, or in labelling or in testing, or in documentation of any of the drug pertaining to this Registration Certificate. In such cases, where there shall be any major change/modification in manufacturing, or in processing or in testing, or in documentation as the case may be, at the discretion of the licensing authority, the manufacturer or his authorised agent in India shall obtain necessary approval within 30 days by submitting a separate application along with the registration fee, as specified in clause (ii) of sub rule (3) of rule 24-A.
6. The manufacturer or his authorised agent in India shall inform the licensing authority immediately in writing in the event of any change in the constitution of the firm and /or address of the registered office/ factory premises operating under this Registration Certificate. Where any such change in the constitution of the firm and/or address takes place, the current Registration Certificate shall be deemed to be valid for a maximum period of three months from the date on which the change has taken place unless, in the meantime, a fresh Registration Certificate has been taken from the licensing authority in the name of the firm with the changed constitution of the firm and/or changed address of the registered office or factory premises".



GOVERNMENT OF INDIA

Central Drugs Standard Control Organisation  
Ministry of Health & Family Welfare

FDA BHAWAN, NEW DELHI (INDIA)

NAME (S) OF DRUGS, WHICH MAY BE IMPORTED  
UNDER REGISTRATION CERTIFICATE NO. BD-917  
DATED 28 APR 2011

1. PREDNISOLONE IP
2. PREDNISOLONE ACETATE USP
3. HYDROCORTISONE IP



ITEMS (THREE) ONLY

सत्यमेव जयते

Place: New Delhi

Date: 28 APR 2011



LICENSING AUTHORITY  
 डा. सुरिन्द्र सिंह  
 Dr. SURINDR SINGH  
 औषधि नियंत्रक (भारत) / Drugs Controller (India)  
 स्वास्थ्य सेवा महानिदेशक  
 Dte. General of Health Services  
 FDA Bhawan, Kotla Road,  
 New Delhi-110002



From :  
The Drugs Controller General (India)  
Directorate General of Health Services

FDA Bhawan, New Delhi  
Dated 28 APR 2011

To,

M/S. PFIZER PRODUCTS INDIA PVT. LTD.,  
SHREE ARIHANT COMPOUND,  
KALHER VILLAGE, BLDG. NO. D7,  
GALA NO. 2, 3 & 4, DIST. THANE,  
BHIWANDI 421 302

Sub: -Registration of M/S. ACTAVIS ITALY S.P.A., VIALE PASTEUR 10, 20014,  
NERVIANO (MI), ITALY under the Provisions of Drug & Cosmetics Rules for the  
purpose of import of drugs in India

Dear Sir,

Please refer to your application no. REG/PPIPL/RC/11/MP/036 dated 17/02/2011: received by this Office vide diary No: 8354 dated: 21/02/2011 on the above subject. Fresh Registration Certificate in Form 41 under the Rules due to change in the name of manufacturer is issued for the manufacturing site along with the name (s) of drug (s) imported under the said Certificate subject to the condition:

1. The drugs shall conform to the standards / specifications mentioned in the Second Schedule of the Act, or such other standards / specifications forwarded by you and approved by this Directorate.
2. The drugs shall have the composition, strengths, standards and packing as approved by this Directorate.
3. Dispute, if any, in respect of the payment of fees and submission of TR6 challan, shall have to be settled between the bank and the applicant.
4. The drugs will be required to be withdrawn from sale from the market in case any undesirable reactions due to its medication are brought to light at any stage. This Directorate should be informed of adverse reports on the drug, if any.
5. The drugs shall be marketed for indications for which they have been approved.
6. If any claim is made for these drugs in future other than those approved, prior permissions from this Directorate will be necessary before such claim is made.
7. The drugs shall be supplied and marketed as per the provisions of the existing Drugs & Cosmetics Act and Rules thereunder.

- 8 This registration in no way relieves you of the responsibility of complying with all other provisions of the Drugs & Cosmetics Act and Rules there under, and any other provisions of any other Act and Rules applicable in the matter concerned.
- 9 Each consignment of the drugs to be imported by you shall be accompanied by a test / analysis reports.
- 10 Based on this registration, import applications for the drug (s) endorsed in the enclosed Registration Certificate shall be considered for the issue of Form 10 Licence under the Rules.
- 11 This Registration Certificate is being issued under the condition that during the pendency of Registration, the applicant may be required to deposit inspection fee as stipulated under Clause (5) of Rule 24-A and enable inspection of manufacturing site by the officials authorised for this purpose. Non-compliance to this condition as and when so directed would result in cancellation of Registration Certification.
- 12 Copy of revalidated Drug Sale License shall be submitted to this office before import.
- 13 Original labels and package inserts of the products as registered by this office vide the Registration Certificate shall be submitted to this office before import.

Please note that Registration Certificate issued is liable to be suspended / cancelled, if any of the condition stipulated above is not complied with, apart from any other condition that may be taken under the provisions of the Drugs & Cosmetics Act, 1940 and the Rules there under.

Please acknowledge the receipt.

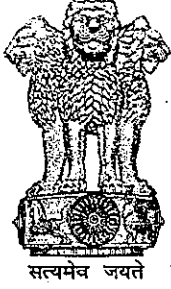
Yours faithfully,

  
Drugs Controller General (India)

**Copy forwarded to:**

1. The Assistant Drug Controller (I), New Custom House, Annexe, Ballard Estate, Fort, Mumbai-400038.
2. The Assistant Drug Controller (I), 15/1, Strand Road, Custom House, Kolkata-700001.
3. The Assistant Drug Controller (I), Room No. 66, 2<sup>nd</sup> Floor, Custom House, Chennai-600001.
4. The Assistant Drug Controller (I), Jawaharlal Nehru Port, Trust Nhava Seva Port, Raigard, Maharashtra.
5. The Assistant Drug Controller (I), International Air Cargo Complex, Indira Gandhi International Air Port, New Delhi-110037.

# GOVERNMENT OF INDIA



## Central Drugs Standard Control Organisation Ministry of Health & Family Welfare

FDA BHAWAN, NEW DELHI (INDIA)

Form 41

(See Rule 27-A)

### REGISTRATION CERTIFICATE

Registration Certificate issued for import of drugs into India

Under Drugs and Cosmetics Rules, 1945

28 APR 2011

Dated

Registration Certificate No. FF-579

1. M/S. ACTAVIS ITALY S.P.A., ITALY having factory premises at M/s. ACTAVIS ITALY S.P.A., VIALE PASTEUR 10, 20014, NERVIANO (MI), ITALY has been registered under Rule 27-A as a manufacturer and is hereby issued this Registration Certificate.
2. Name(s) of drug(s) which may be imported under this Registration Certificate (Please refer the enclosed list of drugs).
3. This Registration Certificate shall be in force from 01/04/2011 to 31/03/2014 unless it is sooner suspended or cancelled under the rules.
4. This Registration Certificate is issued through the office of the manufacturer or his authorised agent in India M/S. PFIZER PRODUCTS INDIA PVT. LTD., SHREE ARIHANT COMPOUND, KALHER VILLAGE, BLDG. NO. D7, GALA NO. 2, 3 & 4, DIST. THANE, BHIWANDI 421 302 who will be responsible for the business activities of the manufacturer in India in all respects.
5. This Registration Certificate is subject to the conditions stated overleaf and to such other conditions as may be specified in the Act and the Rules, from time to time.

Place: New Delhi

Date: 28 APR 2011

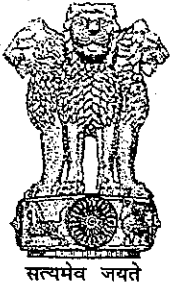


LICENSING AUTHORITY

डा. सुरिंदर सिंग  
Dr. Surinder Singh  
औद्योगिक सेवा (भारत) / Drugs Controller (India)  
औद्योगिक सेवा महानिदेशालय  
Dir. General of Health Services  
FDA Bhawan, Kolda Road,  
New Delhi-110032

Conditions of the Registration Certificate

1. The Registration Certificate shall be displayed at a prominent place by the authorised agent.
2. No drug shall be registered unless it has a free sale approval in the country of origin, and/or in other major countries.
3. The manufacturer or his authorised agent in India shall comply with the conditions of the import licence issued under the Drugs and Cosmetics Rules, 1945.
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5. The manufacturer or his authorised agent in India shall inform the licensing authority within 30 days in writing in the event of any change in manufacturing process, or in packaging, or in labelling or in testing, or in documentation of any of the drug pertaining to this Registration Certificate. In such cases, where there shall be any major change/modification in manufacturing, or in processing or in testing, or in documentation as the case may be, at the discretion of the licensing authority, the manufacturer or his authorised agent in India shall obtain necessary approval within 30 days by submitting a separate application along with the registration fee, as specified in clause (ii) of sub rule (3) of rule 24-A.
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सत्यमेव जयते

GOVERNMENT OF INDIA  
Central Drugs Standard Control Organisation  
Ministry of Health & Family Welfare

FDA BHAWAN, NEW DELHI (INDIA)

NAME (S) OF DRUGS, WHICH MAY BE IMPORTED  
UNDER REGISTRATION CERTIFICATE NO. FF-579  
DATED 28 APR 2011

1. IDARUBICIN HYDROCHLORIDE INJ. USP 5mg, 10mg & 20 mg  
(ZAVEDOS)
2. DAUNORUBICIN HYDROCHLORIDE INJ. USP 20mg  
(DAUNOMYCIN)
3. EPIRUBICIN HYDROCHLORIDE INJ. 10mg & 50mg  
(FARMORUBICIN RD)
4. DOXORUBICIN HYDROCHLORIDE INJ. IP 10 mg & 50mg  
(ADRIAMYCIN RD)
5. IDARUBICIN HYDROCHLORIDE CAPSULES 5mg, 10mg & 25mg  
(ZAVEDOS)

ITEMS (FIVE) ONLY

Place: New Delhi

Date: 28 APR 2011



LICENSING AUTHORITY

Dr. SURINDER SINGH  
श्री. सुरिंदर सिंह (भारत) / Drugs Control Officer  
आरोग्य सेवा महानिदेशक  
Dy. General of Health Services  
FDA Bhawan, Kotla Road,  
New Delhi-110002

