



GOVERNMENT OF THE PEOPLE'S REPUBLIC OF BANGLADESH
MINISTRY OF HEALTH & FAMILY WELFARE
DIRECTORATE OF DRUG ADMINISTRATION
105-106 MOTIJHEEL COMMERCIAL AREA,
DHAKA-1000, BANGLADESH.

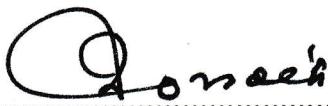
CERTIFICATE OF REGISTRATION

We hereby declare that **PNEUMOVAX 23 (PNEUMOCOCCAL VACCINE POLYVALENT)** manufactured by **Merck Sharp & Dohme Corp. 770 Sumneytown Pike P.O. Box 4, West Point, Pennsylvania, 19486-0004 USA** represented by **Healthcare Pharmaceuticals Ltd, 3/34 East Rampura, Dhaka-1219, Bangladesh** is registered with Directorate of Drug Administration & Licensing Authority (Drugs) under Registration No. **213-4860-14**..... The drug as described below is allowed to be imported into Bangladesh under **The Drugs Act, 1940 (XXIII of 1940)** and **The Drug (Control) Ordinance, 1982** subject to the provision of import Policy published by the Government from time to time.

Name of Product : PNEUMOVAX 23 (PNEUMOCOCCAL VACCINE POLYVALENT)
Dosage Form : Single dose vial
Pack Size : 0.5ml X 1 Dose Vial
Composition : Each 0.5ml Dose of Vaccine Contains 25mcg of Each Polysaccharide Type

<u>Active Ingredients</u>	<u>Specification</u>	<u>Quantity</u>
Pneumococcal Polysaccharide Types 1, 2, 3, 4, 5, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15B, 17F, 18C, 19F, 19A, 20, 22F, 23F, 33F.	INN	25mcg/dose of each type
Other ingredient(s): Sodium Chloride Liquefied Phenol Water for Injections	Ph. Eur. USP Ph. Eur	0.9 % 0.25 % as Phenol QS

Instructions
To be dispensed only by or
on the prescription of a
registered physician.


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Maj. Gen. Md. Jahangir Hossain, **27 APR 2014**
Director General
Directorate of Drugs Administration
&
Licensing Authority (Drugs)
Government of The People's Republic of Bangladesh

Conditions: (i) Labeling should be done in accordance with the provision of the Drugs Act and Rules which require that the name and address of the Manufacturer, Batch Number, Manufacturing date, expiry date, M.R.P (Maximum Retail Price), DAR No. (Drug Administration Registration Number) etc. should be displayed on the label or container and also on the outer cover containing the container. (ii) The registration will be valid for 5 (Five) years from its date of issue unless it is revoked, suspended or cancelled earlier. (iii) The Certificate will be treated as cancelled in any violation of the conditions and if the name or formula of this product changed or modified qualitatively or quantitatively without due approval of the Licensing Authority.

Memo No. **DA/15-5/2010(213)/ 5452**

Dated: **27-04-14**

Copy to: **Healthcare Pharmaceuticals Ltd,**
JANATA TRADERS Principal office:
62/2, Purana Paltan, 3/34 East Rampura, Dhaka-1219
Dhaka-1000, Bangladesh


27.04.14
For Director General
Directorate of Drugs Administration

This Registration shall remain valid
up to 27-04-2024 unless it is suspended
Cancelled or revoked earlier.

Malahuddin
13.01.2020

Md. Salahuddin
Assistant Director
For Director General
Directorate General of Drug Administration
&
Licensing Authority (Drugs)
Government of the People's Republic of Bangladesh



To be dispensed only by the
registered pharmacist
on the prescription of a
physician