

<b>Mebendazole (Mebendazole)</b>			
<b>No. : RPB 203</b>	<b>Rev : 02</b>	<b>Eff. date : 18 07 10</b>	<b>Sign : /</b>

**1 Quality**

- 1.1 USP 37, 2014
- 1.2 Farmakope Indonesia V, 2014
- 1.3 Inhouse Indofarma

**2 Additional Requirements**

- 2.1 Arrival Date : NMT. 12 months from manufacturing date

**3 Certificate of Analysis from Manufacturer**

To be stated on the certificate

- 3.1 Manufacturer name and address
- 3.2 Product name, batch/lot number
- 3.3 Results of test and assays
- 3.4 Compliance with Quality
- 3.5 Quality designation
- 3.6 Date of manufacturing, expiration (year-month) and date of released (year- month-day)
- 3.7 Approval/signature of QA/QC responsibility

**4 Package**

Cardboard drum @ 25 kg, with original seal (inside and outside) from manufacture.

To be stated on goods , label original from manufacture at primary and secondary package each. Preserve in well closed containers.

**5 Labeling**

To be stated on goods **label original from manufacturer**, packing list and invoice :

- 5.1 Name of manufacturer and address
- 5.2 Product name, batch/lot number, manufacturing and expiration date
- 5.3 Quality designation
- 5.4 Gross/net weight
- 5.5 Number of packages

**6 Reference Documents**

- 6.1 USP 37, 2014
- 6.2 Farmakope Indonesia V, 2014
- 6.3 Inhouse Indofarma

**7 History**

Purchasing specification is revision of RPB 203 revision 01 with change in the quality and deleting the review point.

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**8 Distribution**

In general, a copy of this specification is distributed to :

- 8.1 Procurement Dept.
- 8.2 Quality Control Dept.

**9 Approval**

Preparation steps	Position	Code	Signature	Date
Prepared by	Starting and Packaging Material Testing Assistant Manager	AM	[Signature]	180710
Checked by	Analytical Method Assistant Manager	LB	[Signature]	180710
Approved by	Quality Control Manager	AM	<sup>an.</sup> [Signature]	180710
	Research & Development Manager	LB	[Signature]	180710
	Quality Assurance Manager	PM	[Signature]	180710