


Guidelines for Good Manufacturing Practice

In the "Drug Ordinance 1982" section 15 is mentioned that "Every manufacturer of drugs shall follow the good practices in the manufacture and quality control of drugs recommended by the World Health Organization (WHO) and If any manufacturer does not follow such good practices his manufactured license may be cancelled or suspended." So, the all good practice guideline recommended by WHO is the official guideline for Bangladesh.

This circular is to adopt the WHO Guidelines for all good practice mentioned below:

1. Quality System requirements for National Good Manufacturing Inspectorates: WHO Technical Report Series No. 902, 2002 (Annex-8)
2. Good Manufacturing Practice (GMP): WHO Technical Report Series No. 986, 2014 (Annex -2)
3. Good Laboratory Practice (GLP): WHO Technical Report Series No. 957, 2010 (Annex -1)
4. Good Distribution Practice (GDP): WHO Technical Report Series No. 957, 2010 (Annex -5)
5. Good Cold Chain Management Practices (GCCMP): WHO Technical Report Series No. 961, 2011 (Annex -9)
6. Good Storage Practice (GSP): WHO Technical Report Series No. 908, 2003 (Annex -9)
7. Good Documentation Practice (GDocP): WHO Technical Report Series No. 996, 2016 (Annex -5)
- ✓ 8. Quality Risk Management Practice : WHO Technical Report Series No. 981, 2013

The manufacturer and importer of Bangladesh should comply with the principles set forth in the above guidelines.


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&
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