

## Name and Address of Manufacturer

Medbar Tıbbi Malzemeler Turizm San. ve Ticaret A.Ş.  
1142 sok. No:35 Fatih Mah. Sarnıç - İzmir / Turkey  
Phone: +90 232 2816003 Fax: +90 232 2816648

## DECLARATION OF CONFORMITY

I, Ömer Baran, hereby declare that the below mentioned medical device-

- (i) complies with all the requirements under the Act;
- (ii) has been classified according to the classification rules as specified in First Schedule on Rules of Classification of Medical Device; and
- (iii) conforms to requirements specified in APPENDIX 1 of Third Schedule on Essential Principles for Safety and Performance of Medical Devices under Medical Devices Regulations 2012.

## (A) Particulars of medical device

**Generic name:** Emesis Bag

**Specified name:** Emesis Bag (Ref. No:228 02)

**Brand/model:** Medbar

**Manufacturer:** Medbar Tıbbi Malzemeler Turizm San. ve Ticaret A.Ş.

**Country of origin:** TURKEY

**Manufacturing site:** Medbar Tıbbi Malzemeler Turizm San. ve Ticaret A.Ş, 1142 sok. No:35 İzmir / Turkey

**Risk-based classification:** Class I other (not sterile and measuring function )

**Classification rule:** Rule 1

**GMDN code:** 36258

## (B) EN ISO 13485:2012 Quality Management System certificate ("QMS")

**Conformity Assessment Body issuing the certificate:** Trans Pacific Certification Limited

**Certificate number:** MD-0884

**Issuance date:** Feb. 24, 2017

**Expiry date:** Feb. 23, 2017

## (C) Standards Applied

- EN ISO 13485: Medical devices - Quality management systems - Requirements for regulatory purposes
- EN ISO 14971: Medical devices - Application of risk management to medical device
- EN ISO 10993-1: Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process
- EN ISO 15223-1: Medical devices -- Symbols to be used with medical device labels, labelling and information to be supplied -- Part 1: General requirements
- TS EN 1041+A1: Information supplied by the manufacturer of medical devices

I am fully responsible with all the information provided in this declaration. This declaration of conformity is valid from 05. December.2017.

I fully understand and acknowledge that it is an offence under Section 76 of the Medical device Act 2012 [Act 737] to make, sign or furnish any declaration, certificate or other document which is untrue, inaccurate or misleading.

Authorised Signatory:



**medbar**  
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Ömer Baran/General Manager ,05.12.2017

