AYSET TIBBI URUNLER VE PLASTIK TEKSTIL ELEKTRONIK GIDA TEMIZLIK MADDELERI SAN. A.. Sarihamzali Koyu Yolu Uzeri No:54 Seyhan Adana, Turkey Caihan International Marketing whose registered office is at Kotterner Str. 46 D8-7435 Kempten, GERMANY

D15-06

European Declaration of Conformity to the In Vitro Diagnostic Medical Devices, 98/79/EC



Product Name VACCUM TUBE

Product Model Number(s) with K3EDTA, 0,25 ml of blood, sterile
Lot Number 10894-7 / Expire date: 2019.10

DescriptionBlood tube is a device to collect blood samples from the body for

eventual laboratory testing.

Sterile Yes

Classification / Rule Others, According to the Appendix VII

Conformity Assessment Route Annex II of IVD 98/791EC

Applicable Standards EN 980:2003 EN ISO 14971, EN 591:2001 EN ISO 13485:2003

Declaration

AYSET TIBBI URUNLER VE PLASTIK TEKSTIL ELEKTRONIK GIDA TEMIZLIK MADDELERI SAN. A.S. declares that the above product to which this declaration relates, bears the CE Marking, and is in conformity with the applicable requirements of the Directive 98179/EC of the European Parliament and of the Council of 27 October 1998 on in vitro diagnostic medical devices, which allows its free distribution, sale and circulation in EEC.

All supporting documentation is retained at the manufacturer's premises

The present EC Declaration of Conformity is applicable to all mentioned in vitro diagnostic medical devices, manufactured by AYSET TIBURUNLER VE PLASTIK TEKSTIL ELEKTRONIK GIDA TEMIZLIK MADDELERI SAN. A.S. Adana, Turkey, and/or anyway realized under the Manufacturer's Certified Quality System control for a period of about 1 year from the approval date of the present document.

Start date of CE Mark : 21.11.2016

Signature on behalf of

AYSETTIBBI URUNLER VE PLASTIKTEKSTIL ELEKTRONIK GIDATEMIZLIK MADDELERI SAN.AS.

Date Signature 2 1.11.2016

HASAN AYTEKIN

AYSET General MANAGER