## Notification of a Body in the framework of a technical harmonization directive

From:

Ministriy of Trade – DG Product Safety and Inspection Söğütözü Mah. 2176. Sk. No:63 06530 Çankaya Ankara Türkiye

To:

European Commission GROWTH Directorate-General 200 Rue de la Loi, B-1049 Brussels.

**Other Member States** 

#### **Reference:**

Legislation: 93/42/EEC Medical devices

#### Body name, address, telephone, fax, email, website :

Notice Belgelendirme, Muayene ve Denetim Hizmetleri Anonim Şirketi Esentepe Mahallesi Milangaz Caddesi No:75 A/92 Kartal/İstanbul Istanbul Türkiye +90 (216) 504 16 98 info@notice.com.tr www.notice.com.tr

# **Body info:**

NB 2764

**Body:** 

**Created:** 2019-01-03

**Last update:** 2023-11-03

### Name of National Accreditation Body (NAB):

TURKAK (Turkish Accreditation Agency)

The accreditation covers the product categories and conformity assessment procedures concerned by this notification: Yes

Tasks performed by the Body:

Last approval date: 2020-11-19

| Products  | Procedures  | Articles /Annexes   | Limitations                               |
|---|---|---------------------|---|
|   |   |                     |   |
| -*MD 0100 - General non-<br>active, non-implantable<br>medical devices<br>-*MD 0101 - Non-active<br>devices for anaesthesia,<br>emergency and intensive<br>care     | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0100 - General non-<br>active, non-implantable<br>medical devices<br>-*MD 0102 - Non-active<br>devices for injection,<br>infusion, transfusion and<br>dialysis | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0100 - General non-<br>active, non-implantable<br>medical devices<br>-*MD 0104 - Non-active<br>medical devices with<br>measuring function                      | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0100 - General non-<br>active, non-implantable<br>medical devices<br>-*MD 0105 - Non-active<br>ophthalmologic devices  | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0100 - General non-<br>active, non-implantable<br>medical devices<br>-*MD 0108 - Non-active<br>medical devices for<br>disinfecting, cleaning, rinsing          | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0200 - Non-active<br>implants<br>-*MD 0201 - Non-active<br>cardiovascular implants   | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex V             | Only felts and<br>similar<br>technologies |
| -*MD 0200 - Non-active<br>implants<br>-*MD 0202 - Non-active<br>orthopaedic implants  | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0200 - Non-active<br>implants<br>-*MD 0203 - Non-active<br>functional implants   | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0200 - Non-active<br>implants<br>-*MD 0204 - Non-active soft<br>tissue implants  | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |
| -*MD 0300 - Devices for<br>wound care<br>-*MD 0301 - Bandages and<br>wound dressings  | EC declaration of conformity (full<br>quality assurance system)<br>EC declaration of conformity<br>(production quality assurance) | Annex II<br>Annex V |   |

| Products                                  | Procedures   | Articles /Annexes   | Limitations    |
|---|--|---------------------|----------------|
| FIGURES                                   | Frocedures   | Articles /Armexes   |                |
| -*MD 0300 - Devices for                   | EC declaration of conformity (full                           | Annex II            |                |
| wound care                                | quality assurance system)                                    | Annex V             |                |
| -*MD 0302 - Suture material<br>and clamps | EC declaration of conformity (production quality assurance)  |                     |                |
| and clamps                                |  |                     |                |
| -*MD 0300 - Devices for                   | EC declaration of conformity (full                           | Annex II            |                |
| wound care                                | quality assurance system)                                    | Annex V             |                |
| -*MD 0303 - Other medical                 | EC declaration of conformity                                 |                     |                |
| devices for wound care                    | (production quality assurance)                               |                     |                |
| -*MD 0400 - Non-active                    | EC declaration of conformity (full                           | Annex II            |                |
| dental devices and                        | quality assurance system)                                    | Annex V             |                |
| accessories                               | EC declaration of conformity                                 |                     |                |
| -*MD 0401 - Non-active                    | (production quality assurance)                               |                     |                |
| dental equipment and                      |  |                     |                |
| instruments                               |  |                     |                |
| -*MD 0400 - Non-active                    | EC declaration of conformity (full                           | Annex II            |                |
| dental devices and                        | quality assurance system)                                    | Annex V             |                |
| accessories                               | EC declaration of conformity                                 |                     |                |
| -*MD 0402 - Dental materials              | (production quality assurance)                               |                     |                |
| -*MD 0400 - Non-active                    | EC declaration of conformity (full                           | Annex II            |                |
| dental devices and                        | quality assurance system)                                    | Annex V             |                |
| accessories                               | EC declaration of conformity                                 |                     |                |
| -*MD 0403 - Dental implants               | (production quality assurance)                               |                     |                |
| -*MD 1100 - General active                | EC declaration of conformity (full                           | Annex II            | Only infusion  |
| medical devices                           | quality assurance system)                                    |                     | devices        |
| -*MD 1101 - Devices for                   | EC declaration of conformity                                 |                     |                |
| extra-corporal circulation,               | (production quality assurance)                               |                     |                |
| infusion and haemopheresis                |  |                     |                |
| -*MD 1100 - General active                | EC declaration of conformity (full                           | Annex II            | excluding      |
| medical devices                           | quality assurance system)                                    | Annex V             | hyperbaric     |
| -*MD 1102 - Respiratory                   | EC declaration of conformity                                 |                     | chambers for   |
| devices, devices including                | (production quality assurance)                               |                     | oxygen therapy |
| hyperbaric chambers for                   |  |                     |                |
| oxygen therapy, inhalation<br>anaesthesia |  |                     |                |
|   |  |                     |                |
| -*MD 1100 - General active                | EC declaration of conformity (full                           | Annex II            |                |
| medical devices                           | quality assurance system)                                    | Annex V             |                |
| -*MD 1104 - Active surgical               | EC declaration of conformity                                 |                     |                |
| devices                                   | (production quality assurance)                               |                     |                |
| -*MD 1100 - General active                | EC declaration of conformity (full                           | Annex II            |                |
| medical devices                           | quality assurance system)                                    | Annex V             |                |
| -*MD 1111 - Software                      | EC declaration of conformity                                 |                     |                |
| *MD 1200 Manitarian                       | (production quality assurance)                               | Appoy               |                |
| -*MD 1300 - Monitoring<br>devices         | EC declaration of conformity (full quality assurance system) | Annex II<br>Annex V |                |
| -*MD 1301 - Monitoring                    | EC declaration of conformity                                 |                     |                |
| devices of non-vital                      | (production quality assurance)                               |                     |                |
| physiological parameters                  |  |                     |                |
| -*MD 1300 - Monitoring                    | EC declaration of conformity (full                           | Annex II            |                |
| devices                                   | quality assurance system)                                    | Annex V             |                |
| -*MD 1302 - Monitoring                    | EC declaration of conformity                                 |                     |                |
| devices of vital physiological            | (production quality assurance)                               |                     |                |
| parameters                                |  |                     |                |
|   |  |                     |                |

| Products                     | Procedures  | Articles /Annexes   | Limitations |
|------------------------------|---|---------------------|-------------|
| radiation therapy and thermo | quality assurance system)<br>EC declaration of conformity | Annex II<br>Annex V |             |

| Horizontal technical competences  | Limitations  |
|---|--|
| *MDS 7001 - Medical devices incorporating<br>medicinal substances, according to Directive<br>2001/83/EC:                    |  |
| *MDS 7004 - Medical devices referencing the<br>Directive 2006/42/EC on machinery:   |  |
| *MDS 7006 - Medical devices in sterile condition:   | Including aseptic, processing, ethylene oxide gas,<br>sterilisation (EOG), moist heat sterilisation,<br>radiation sterilisation (gamma, x-ray, electron<br>beam) |
| *MDS 7009 - Medical devices utilising biological<br>active coatings and/or materials or being wholly<br>or mainly absorbed: |  |
| *MDS 7010 - Medical devices incorporating<br>software /utilising software /controlled by<br>software:                       |  |