

**DEPARTMENT OF HEALTH**  
**Directorate: Radiation Control**

**APPLICATION FOR A LICENCE TO IMPORT  
A NEW LISTED ELECTROMEDICAL DEVICE**

HAZARDOUS SUBSTANCES ACT, 1973 (ACT 15 OF 1973)

Postal Address: Directorate: Radiation Control, Private Bag X62, Bellville, 7535  
Street Address: 2nd Floor, Louwville Place, cor. Vrede & Kort St., Bellville, 7530

Enquiries: Tel: 021 - 948 6162 Fax: 021 - 946 1589

**A: PRIMARY IMPORTER - APPLICANT**

|                        |                  |                        |
|------------------------|------------------|------------------------|
| <b>Name:</b>           |                  |                        |
| <b>Postal Address:</b> |                  | <b>Street Address:</b> |
|                        |                  |                        |
|                        |                  |                        |
|                        | <b>Postcode:</b> |                        |
| <b>Website:</b>        |                  |                        |

**B: PRODUCT INFORMATION**

|  |
|--|
| <b>Brand:</b>  |
| <b>Model:</b>  |
| <b>Intended purpose of this device according to the manufacturer's labelling and instructions for use:</b> |
|  |
|  |

**C: MANUFACTURER**

|                 |
|-----------------|
| <b>Name:</b>    |
| <b>Address:</b> |
|                 |
| <b>Website:</b> |

**D: AUTHORISED REPRESENTATIVE IN THE EUROPEAN UNION**

|                 |             |
|-----------------|-------------|
| <b>Name:</b>    |             |
| <b>Address:</b> |             |
|                 |             |
| <b>Website:</b> |             |
| <b>Email:</b>   | <b>Fax:</b> |

**E: COMPANY CONTACT PERSON (for all regulatory correspondence)**

|   |              |
|---|--------------|
| I, ..... declare all the information supplied to be correct and true. |              |
| Signature:  | Date:        |
| Title (Mr, Ms, Mrs, Dr, etc.):  | Designation: |
| Tel:  | Cell:        |
| Email:  | Fax:         |

**REQUIREMENTS**

re

**APPLICATION FOR A LICENCE TO IMPORT  
A NEW LISTED ELECTROMEDICAL DEVICE**

The applicant must supply the following documentation for **each** model to be imported:

(Please note: \* Faxed applications will not be acceptable;

\* The electronic version of any document will be acceptable only if it is in either MS Word or pdf format)

- Annexure 1: Completed application form 41BM-1(IMP); *and*
- Annexure 2: Colour brochure (including technical specifications); *and*
- Annexure 3: Letter of appointment as the sole authorised agent/representative of the original manufacturer in South Africa; *and*
- Annexure 4: **EC Certificate(s) issued by a Notified Body** in terms of EC Directive 93/42/EEC or 90/385/EEC (whichever one is applicable); *and*
- Annexure 5: **EC Declaration of Conformity by the manufacturer** in terms of EC Directive 93/42/EEC or 90/385/EEC (whichever one is applicable).