



Project and Product Certification Representative Authorization Letter

| Reason for Amendment (current / obsolete) | Revision History | | |
|---|------------------|-----|---------------|
| | From | То | Approved Date |
| Initial Release (obsolete) | 1.0 | 1.0 | Nov-14-2006 |
| Revised wording (obsolete) | 1.0 | 2.0 | Sept-25-2007 |
| Updated company template (obsolete) | 2.0 | 3.0 | Jan-31-2012 |
| Updated letter information (obsolete) | 3.0 | 4.0 | May-23-2014 |
| Added FCC ID field (current) | 4.0 | 5.0 | Sept 16 2014 |

Medtronic

3/27/2018

To: SIEMIC, Inc. 775 Montague Expressway, Milpitas, CA 95035 USA

Dear Sir/Madam,

Re: Project and Product Certification Representative Authorization Letter

We, <u>Medtronic</u>, <u>Inc.</u> hereby authorize SIEMIC, Inc. to act as Certification Body for certifying for the following project(s):

Product: MRI rechargeable neurostimulator - InterStim Micro

FCC ID: LF597810

We affirm that between SIEMIC, Inc. and <u>Medtronic, Inc.</u>, any difference in understanding, including test plan, measurement methods, applicable standards and relevant procedures and processes have been resolved prior to commencement of testing activities.

Sincerely,

Guitaume Girard

EMC and RF Regulatory Compliance

Technical Fellow Medtronic, Inc. Tel 763-526-0652

g.guillaume@medtronic.com