

Activa® SC Multi-program neurostimulator 37603

Implant manual

!USA Rx only



2010

Explanation of symbols on product or package labeling

Refer to the appropriate product for symbols that apply.



Open here



Do not reuse



Sterilized using ethylene oxide



Caution, consult accompanying documents



Date of manufacture



Manufacturer



Use by



Temperature limitation



Serial number



Conformité Européenne (European Conformity). This symbol means that the device fully complies with AIMD Directive 90/385/EEC (NB 0123) and R&TTE Directive 1999/5/EC.



Authorized representative in the European community



For USA audiences only

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Refer to the Indications sheet for indications and related information.

Refer to the Information for Prescribers booklet for contraindications, warnings, precautions, adverse events summary, patient selection, resterilization, and component disposal.

Refer to the System Eligibility Battery Longevity reference manual for neurostimulator selection and battery longevity calculations.

I-USA Refer to the Clinical Summary booklet for information on the clinical study results of the neurostimulation system, individualization of treatment, and use in specific populations.

Description

The Medtronic Activa SC Model 37603 Neurostimulator is part of a neurostimulation system for deep brain stimulation.

Package contents

- Neurostimulator
- Torque wrench
- Product literature
- !USA Warranty card
- Registration form
- Patient identification card

Patient identification card

A patient identification card is packaged with this device. Advise the patient to carry the identification card at all times

! USA The patient identification card packaged with the device is temporary; a permanent card will be mailed to the patient when Medtronic receives the registration form.

The implant registration form registers the device warranties and creates a record of the device in Medtronic's implant data system.

Device specifications

The neurostimulator is a multi-program device that delivers stimulation through one lead. The stimulation settings are stored in programs. A program is a specific combination of pulse width, rate, and amplitude settings acting on a specific electrode combination. Up to two programs can be combined into a group. When using more than one program, the pulses are delivered sequentially—first a pulse from one program, then a pulse from the next program.

Pulse width, amplitude, and electrode polarity for each program within the group can have different values. Rate, rate limits, SoftStart/Stop and Cycling for each program within the group have the same values

Table 1. Operating values for the Activa SC Model 37603 neurostimulator

Programmable parameter	Operating range and resolution ^a
Number of defined groups	1 to 4
Number of programs per group	1 to 2
Electrode configuration ^b	1 to 4 electrodes per lead as anode (+), cathode (-), or OFF; case defined as anode or OFF
Amplitude (voltage mode)	0 to 10.5 V with 0.05-V or 0.1-V resolution
Amplitude (current mode)	0 to 25.5 mA with 0.1-mA resolution
Amplitude – upper patient limit	Tracking limit (by hemisphere): +0 to +2 (0.2 resolution); +2 to +4.5 (0.5 resolution)
Amplitude – lower patient limit	Tracking limit (by hemisphere): -0 to -2 (0.2 resolution); -2 to -4.0 (0.5 resolution); full range $^{\rm c}$
Pulse width	60 to 450 μs (10-μs resolution)
Pulse width – upper patient limit	Tracking limit: +0 to +100 μs (10-μs resolution)
Pulse width – lower patient limit	Tracking limit: -0 to -100 μs (10-μs resolution)
Rate (voltage mode)	2 to 250 Hz (resolution: 1 Hz from 2 Hz to 10 Hz, 5 Hz from 10 Hz to 250 Hz) $^{\rm d}$
Rate (current mode)	30 to 250 Hz (5-Hz resolution) ^d
Rate – upper patient limit	Tracking limit: +0 to +50 Hz (10-Hz resolution)
Rate – lower patient limit	Tracking limit: -0 to -50 Hz (10-Hz resolution)
SoftStart/Stop	OFF, ON: 1-, 2-, 4-, or 8-second ramp duration
Cycling	OFF, ON: 0.1 s to 24 hr (resolution: 0.1 s from 0.1 s to 1 s, 1 s from 1 s to 59 s, 1 min from 1 min to 59 min, 1 hr from 1 hr to 24 hr)

^a Interlocks will prevent the use of some parameter combinations.

^b In current mode, a maximum of 2 electrodes (including the case) can be configured: one as anode (+) and one as cathode (-).

c Full range = -10.5 V (voltage mode); -25.5 mA (current mode).

d Rate limited to 125 Hz when two programs are active on a single lead.

Table 2. Physical characteristics of the Activa SC Model 37603 neurostimulatora

Description	Value	
Connector type	Octapolar, one bore ^{bc}	
Height	55 mm (2.2 in)	
Length	60 mm (2.4 in)	
Thickness	11 mm (0.4 in)	
Weight	44 g (1.6 oz)	
Volume	27 cm ³	
Power source	4.5 Amp hours, 3.2 V HCSVOd cell	
Temperature limitation	-18° to +52°C (0° to +126°F)	
Serial number model designatore	NLB	
Radiopaque identification (ID) code	NLB	

a All measurements are approximate.

b Compatible with 8-4 extension.

c Four contacts active.

d Hybrid combined silver vanadium oxide.

e The serial number is the model designator followed by a number. The clinician programmer displays the entire serial number beginning with the model designator.

Table 3. Material of components in the Activa SC Model 37603 package

Components	Material	Material contacts human tissue
Neurostimulator		
Case	Titanium, parylene	Yes
Connector block	Siloxane coated polysulfone, silicone medical adhesive	Yes
Grommets, seals	Silicone rubber	Yes
Setscrews	Titanium alloy	Yes
Adhesive	Silicone medical adhesive	Yes
Torque wrench		
Handle	Polyetherimide	Yes
Shaft	Stainless steel	Yes

Declaration of conformity

Medtronic declares that this product is in conformity with the essential requirements of Directive 1999/5/EC on Radio and Telecommunications Terminal Equipment and Directive 90/385/EEC on Active Implantable Medical Devices.

For additional information, contact the appropriate Medtronic representative listed on the inside back cover of this manual.

Instructions for use

Implanting physicians should be experienced in stereotactic and functional neurosurgery and deep brain stimulation procedures, as well as thoroughly familiar with all product labeling.



Cautions:

- When using sharp instruments near the neurostimulator, be extremely careful to avoid nicking or damaging the case, the insulation, or the connector block.
 Damaging the neurostimulator may require surgical replacement.
- Do not use saline or other ionic fluids at connections, which could result in a short circuit.

Verifying neurostimulator operation

Before opening the sterile neurostimulator package, verify that the neurostimulator is operable by using the clinician programmer to interrogate the neurostimulator and read the neurostimulator battery service life. (Refer to the software manual for instructions on how to read the battery service life.)



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 \sum \text{Caution:} \] Do not implant a neurostimulator if it was dropped onto a hard surface from a height of 30 cm (12 in) or more, because the neurostimulator may be damaged and fail to operate properly.

Note: The neurostimulator pocket may be flushed with an antibiotic solution; do not submerge the neurostimulator in fluid.

Connecting the extension to the neurostimulator



Caution: Before connecting components, wipe off any body fluids and dry all connections. Fluids in the connection may result in stimulation at the connection site, intermittent stimulation, or loss of stimulation.

- Wipe the extension connector with sterile gauze. If necessary, use sterile (United States Pharmacopeia [USP]) water or a nonionic antibiotic solution.
- 2. Make sure the connector block receptacle is dry and clean.
- Insert the extension connector into the neurostimulator socket until it is fully seated within the connector block (Figure 1).

Notes:

- During insertion, some resistance is typical.
- To retract the setscrew, insert the torque wrench (packaged with the neurostimulator) into the self-sealing grommet and rotate the setscrew counterclockwise; however, do not remove the setscrew from the connector block.



Caution: Do not insert the extension connector into the connector block if the setscrew is not sufficiently retracted. If the setscrew is not retracted, the setscrew

may damage the extension and the extension will not be seated fully into the connector block.



Figure 1. Insert the extension connector fully into the neurostimulator socket.

For the extension, fully insert the torque wrench into each self-sealing grommet of the connector block and tighten setscrew (Figure 2).



△ Cautions:

- Be sure the torque wrench is fully inserted into the self-sealing grommet. If the torque wrench is not fully inserted, the setscrew may be damaged, resulting in intermittent or loss of stimulation.
- Before tightening the setscrew, ensure that the extension connector pin is inserted into the connector block to prevent damaging the extension.
- Verify that each leaf of the self-sealing grommet is closed after the torque wrench is withdrawn. If fluid leaks through a grommet seal that is not fully closed, the patient may experience shocking, burning, or irritation at the neurostimulator implant location, or intermittent stimulation, or loss of stimulation.



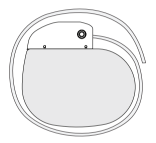
Figure 2. Tightening the setscrew in the self-sealing grommet.

Implanting the neurostimulator

1. Place the neurostimulator into the subcutaneous pocket with the Medtronic logo facing outward, away from muscle tissue, and ensure that the extension is not bent sharply.

Cautions:

- Ensure that the neurostimulator is placed no deeper than 4 cm (1.5 in) below the skin and is parallel to the skin. If the neurostimulator is too deep or is not parallel to the skin, telemetry may be unsuccessful.
- Do not coil excess extension in front of the neurostimulator. Wrap excess extension around the perimeter (Figure 3) of the neurostimulator to minimize subcutaneous pocket depth, help minimize potential damage during neurostimulator replacement surgery, help minimize potential kinking of the extension, and minimize interference with telemetry. Excess extension should not exceed two wraps around the perimeter of the neurostimulator. Extension lengths requiring more than two wraps can interfere with telemetry.



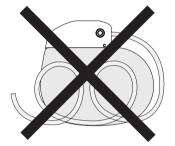


Figure 3. Wrap excess extension around the perimeter of the neurostimulator.

2. Use the suture holes in the connector block to secure the neurostimulator to the muscle fascia with nonabsorbable silk

Checking system integrity



Caution: To use the nonsterile clinician programmer in a sterile field, place a sterile barrier between the patient and the programming head to prevent infection. Do not sterilize any part of the clinician programmer. Sterilization may damage the programmer.

Note: The neurostimulator should be in the pocket during system interrogation for integrity to ensure proper readings.

- To ensure proper connection of the extension to the neurostimulator, use the clinician
 programmer to program the lead configuration and basic stimulation parameters, check
 the battery status, and check the electrode impedances to rule out a short or open circuit.
- 2. If the system integrity test results are not acceptable, refer to "Connecting the extension to the neurostimulator" on page 9.

Completing the implant procedure

- 1. Close and dress all incisions.
- 2. Ensure that a patient control device is given to the patient.
- Complete the device tracking and patient registration paperwork and return the documents to Medtronic.

Contacts:

Asia.

Medtronic International Ltd.

Tel 02891-4068 Fax 02591-0313

Medtronic Asia Ltd.

Tel. (02)-548-1148 Fax (02)-518-4786

Australia:

Medtronic Australasia Pty. Ltd. 97 Waterloo Road

North Rvde, NSW 2113

Australia

Tel. +61-2-9857-9000

Fax +61-2-9878-5100 Toll free 1-800-668-670

Austria:

Medtronic Österreich GmbH

Tel. 01-240440

Fax 01-24044-100

Belaium:

Medtronic Belgium S.A.

Tel. 02-456-0900 Fax 02-460-2667

Canada:

Medtronic of Canada Ltd.

Tel. (1905)-826-6020 Fax (1905)-826-6620

Czech Republic:

Medtronic Czechia s.r.o. Tel. 2-965-795-80

Fax 2-965-795-89

Denmark:

Medtronic Danmark A/S Tel. 45-32-48-18-00

Fax 45-32-48-18-01

Finland:

Medtronic Finland Oy/LTD Tel. (09)-755-2500

Fax (09)-755-25018

France:

Medtronic France S.A.S. Tel. 01-5538-1700

Fax 01-5538-1800

Germanv:

Medtronic GmbH Tel. (02159)-81490

Fax (02159)-8149100

Greece:

Medtronic Hellas S.A. Tel. 210-67-79-099

Fax 210-67-79-399

Hungary:

Medtronic Hungária Kft.

Tel 1-889-06-00

Fax 1-889-06-99

Ireland:

Medtronic Ireland Ltd.

Tel. (01)-890-6522 Fax (01)-890-7220

Italy:

Medtronic Italia SpA

Tel 02-241371

Fax 02-241381

Tel. 06-328141 Fax 06-3215812

Japan:

Medtronic Japan Tel. 3-6430-2011

Fax 3-6430-7140

Latin America:

Medtronic Inc.

Tel. (1305)-500-9328 Fax (1786)-709-4244

Norway:

Medtronic Norge AS

Tel. 067-10-32-00

Fax 067-10-32-10

Poland:

Medtronic Poland Sp. z.o.o.

Tel. (022)-465-69-00

Fax (022)-465-69-17

Portugal:

Medtronic Portugal, Lda.

Tel. 21-724-5100

Fax 21-724-5199

Russia:

Medtronic Russia

Tel. (8495) 580-7377

Fax (8495) 580-7378

Slovakia

Medtronic Slovakia, o.z.

Tel. 0268 206 911

Fax 0268 206 999

Spain: Medtronic Ibérica, S.A.

Tel. 91-625-0400

Fax 91-650-7410

Sweden:

Medtronic AB

Tel. 08-568-585-00

Fax 08-568-585-01

Switzerland:

Medtronic (Schweiz) AG Tel. 031-868-0100 Fax 031-868-0199

The Netherlands:

Medtronic B.V. Tel. (045)-566-8000 Fax (045)-566-8668

U.K.:

Medtronic U.K. Ltd. Tel. 01923-212213 Fax 01923-241004 USA:

Medtronic, Inc. Tel. (1763)-505-5000 Fax (1763)-505-1000 Toll-free: (1-800)-328-0810



Manufacturer (

Medtronic, Inc. 710 Medtronic Parkway Minneapolis, MN 55432-5604 USA

Internet: www.medtronic.com

Tel. 1-763-505-5000 Fax 1-763-505-1000

Medtronic E.C. Authorized EC REP Representative/Distributed by

Medtronic B V Earl Bakkenstraat 10 6422 P.I Heerlen The Netherlands Tel. 31-45-566-8000 Fax 31-45-566-8668

Europe/Africa/Middle East

Headquarters Medtronic International Trading Sàrl Route du Molliau 31 Case Postale 84 CH-1131 Tolochenaz Switzerland Internet: www medtronic com Tel. 41-21-802-7000 Fax 41-21-802-7900

Asia-Pacific

Medtronic International Ltd. Suite 1602 16/F, Manulife Plaza The Lee Gardens, 33 Hysan Avenue Causeway Bay Hong Kong Tel. 852-2891-4068 Fax 852-2591-0313

Contacts for specific countries are listed inside this cover.



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