



PATIENT SYSTEM HANDBOOK

PRECISION Spinal Cord Stimulation System



Patient System Handbook

CAUTION:

Federal law restricts this device to sale, distribution and use by or on the order of a physician.

Copyright

©2006 by Advanced Bionics Corporation. All Rights Reserved. Any copying, reproduction or translation of all or part of the contents of this document without the express written permission of Advanced Bionics Corporation is strictly forbidden by the provisions of the law of March 11th, 1957.

Guarantees

Advanced Bionics Corporation reserves the right to modify, without prior notice, information relating to its products in order to improve their reliability or operating capacity.

Drawings are for illustration purposes only.

Registered Trademarks

Velcro® is a registered mark of Velcro Industries, Manchester, New Hampshire. Other brands and their products are trademarks or registered trademarks of their respective holders and should be noted as such.

Table of Contents

1	Introduction	1
2	System Description	3
3	Safety Information	7
	Indications for Use	
	Precision System Clinical Summary	
	Contraindications	
	Warnings	18
	Precautions	21
	Adverse Effects	26
4	The Remote Control	27
	Buttons and Basic Operation	
	Communicating with the Stimulator	
	Stimulation On and Off	
	Signal Strength	32
	The Level Screen	34
	Stimulation Level Control	34
	Selecting Areas for Level Control	38

	Programs	
	Selecting and Activating Programs	
	Saving Program Changes	42
	Options	45
	Understanding Battery Messages	51
5	Charging the Stimulator	55
	Getting Started	
	Charging Your Stimulator	
6	Help	65
•	Stimulation	
	Remote Control Display Messages	
	Accessories	
	Contacting Advanced Bionics	
7	Limited Warranty	73
	Implanted Pulse Generator	
	Externals	
	Glossary	79
	-	
	Index	

1

Introduction

The Advanced Bionics PrecisionTM SCS (Spinal Cord Stimulation) system is prescribed for the management of chronic pain. The system electrically stimulates the spinal cord to alter the perception of pain signals that move along the nerve pathways on either side of the spine. *Paresthesia* is the term that describes the light, tingling sensation—the "feeling"—of spinal cord stimulation.

Before receiving your new Stimulator, you had the opportunity to test stimulation therapy and decide if it would work for you. By choosing to have a stimulator surgically implanted, you



confirmed that *paresthesia* is capable of providing you with good to excellent pain relief. As you go forward with this therapy, your healthcare professionals will work with you to find the most comfortable level of paresthesia to cover the painful areas by adjusting your Stimulator's settings. Although you may have pain areas that cannot be reached by spinal cord stimulation, the goal is to bring you the most effective pain relief possible. The more you help and work with your healthcare professionals, the more likely you are to achieve the best outcome possible from your new Precision system.

Advanced Bionics is an organization dedicated to helping you manage your pain. We will help you make the most of this therapy for an improved quality of life.

2

System Description

The Precision system includes both implanted and external components. During your surgery, one or more wires called **leads** were placed along your spinal cord where pain signals to the brain can be intercepted. The lead was then attached to an implantable pulse generator (IPG or Stimulator), also referred to as an **implant**, or **Stimulator**. The Stimulator is commonly placed in the abdomen, upper buttock, or subclavicular area. The Stimulator sends a small electrical current to a series of stimulating contacts, or **electrodes**, at the end of the lead. The battery-powered Stimulator is controlled by a hand-held programmer or **Remote Control**, and is periodically recharged using a separate **Charging System**.

The Remote Control, the heart of the Precision system, is a powerful yet easy to use tool for managing every aspect of your pain treatment—from controlling the level, or strength, of stimulation to accessing special treatment programs and program options.

To make the most of your Precision system, it is important to learn:

- 1. what to be aware of for safety,
- 2. how to use the Remote Control, and
- 3. how to recharge the Stimulator.

These subjects are covered on the following pages, and we encourage you to read this manual entirely. If you have any questions, or need clarification of anything contained in this manual, feel free to contact our Customer Service Department at (866) 360-4747.

Before reading more about the Precision System, first check that you have received all of the following items. (And check to be sure you have your Temporary Patient Identification Card; keep it with you until you receive your permanent card.) If any item is missing, please call our Customer Service department for a replacement.

- (1) Remote Control
- (1) Charger
- (1) Charger Base Station
- (1) Power Supply
- (1) Remote Control Case
- (1) Charging Belt
- (1) Charger Adhesives (52 pieces)
- (1) Remote Control Batteries (3)

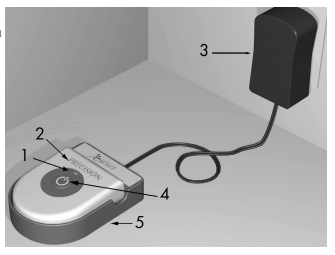
Note: Use only accessories supplied by Advanced Bionics.



Permanent Patient Identification Card

Find a convenient electrical outlet, one that will not expose the Charging System parts to water or direct heat, and plug in the Base Station Power Supply. Next, connect the Power Supply to the Base Station, and locate the Base Station on a flat surface. Finally, place the Charger in the Base Station with the stimulation on/off button facing up.

For now, that is all you need to do to get started. For more information on the Charging System and its use, see "Charging the Stimulator" on page 55.



- 1. Indicator Light
- 2. Charger
- 3. Power Supply

- 4. Power Button
- 5. Base Station

3

Safety Information

Indications for Use

The Advanced Bionics PrecisionTM Spinal Cord Stimulator System (Precision System) is indicated as an aid in the management of chronic intractable pain of the trunk and/or limbs, including unilateral or bilateral pain associated with the following: failed back surgery syndrome, intractable low back pain and leg pain.

Precision System Clinical Summary

Determination of the safety and effectiveness of the PRECISION System was based on available published clinical studies for similar implanted spinal cord stimulation systems. The PRECISION System is similar to the SCS systems reported in published literature in intended use, target patient population, technology, device design, and output characteristics. Therefore, the clinical data from the published literature described below represents evidence supporting the safety and effectiveness of the PRECISION System for

the treatment chronic intractable pain of the trunk and/or limbs, including unilateral or bilateral pain associated with the following: failed back surgery syndrome, intractable low back and leg pain.

Efficacy Evaluation

Three (3) clinical literature studies were used to support the effectiveness of the PRECISION System (Ohnmeiss et al. 1996, Villavincencio et al. 2000, Hassenbach SJ et al. 1995). The studies included a total of 116 patients that were implanted with an SCS system. A total of approximately 3166 device months of experience was depicted from the retrospective clinical evaluation. All three studies examined the effectiveness of SCS on patients with chronic pain of the trunk and/or limbs including unilateral or bilateral pain associated with the following: failed back surgery syndrome or intractable low back and leg pain. In all studies, a totally implantable spinal cord stimulator was used in association with a percutaneous and/or surgical lead. These studies provide the same diagnostic or therapeutic intervention for the same disease/conditions and patient population as the PRECISION System.

The prospective study by Ohnmeiss et al. 1996, examined the long-term effectiveness of SCS in patients with intractable leg pain. Forty patients were implanted with SCS systems and evaluated at 6 weeks, 12 months, and 24 months follow-up. Outcome measures

included the VAS, pain drawings, medication use, SIP (Sickness Impact Profile), isometric lower extremity testing, and patient questionnaires. An intent-to-treat analysis was performed. After patients had SCS for 24 months, leg pain, pain when walking, standing pain, pain's effect on overall lifestyle, and the total analog scale scores were significantly improved from baseline. In this study, 25% of the implanted patients had greater than 50% improvement in pain rating.

In addition, 3 patients from this study had their stimulators repositioned due to pain at the original location. Three patients had reoperations to adjust lead position; 1 patient required 2 reoperations, 1 patient had the device removed due to infection and later to have a new device implanted. A diabetic patient had skin problems which required device removal; a new device was later implanted. Two patients had the device removed due to unsatisfactory pain relief.

The prospective study performed by Villavicencio et al. 2000 included 41 patients with pain of various etiologies. The majority of the patients, 24 (59%), had Failed Back Surgery Syndrome (FBSS), 7 (17%) had Complex Regional Pain Syndrome (CRPS I and II), 4 (10%) had neuropathic pain syndrome, and 6 (15%) were diagnosed as stroke or other. Patients underwent an initial trial period for SCS with temporary leads. If the trial resulted in greater than 50% reduction in the patient's pain, as measured by the VAS, the patient was

implanted with a SCS system. In this study, 27/41 patients, 66%, had permanent implants. All patients were examined after 6 weeks. Pain measurements were assessed at 3-6 month intervals for the first year and annually thereafter. The median long-term follow-up was 34 months. A total of 24/27 (89%), reported greater than 50% reduction in pain. Since the majority of the patients were treated for FBSS, this article supports the use of SCS for the treatment of FBSS.

In this study, one patient required a revision because of electrode fracture. One patient required removal of the system due to local infection. One patient required replacement of the Stimulator due to mechanical failure. Overall, 16 of 27 (59%) patients required a total of 36 repositioning procedures.

A retrospective analysis performed by Hassenbusch SJ et al. 1995 included patients with chronic lower body pain, predominately neuropathic pain and pain either midline lower back and/or unilateral or bilateral leg pain treated over a 5 year period. The study was a comparison of SCS to spinal infusion of opioids. For patients with radicular pain involving one leg with or without unilateral buttock pain, a trial of SCS was recommended first. For patients with midline back pain and /or bilateral leg pain, a trial of long-term spinal infusion was recommended first. If the patients failed screening with either of these modalities, the other was then tested. If the treatment reduced the pain by 50%, the systems were

internalized. A retrospective analysis of patients with unilateral leg and/or buttock pain treated initially with SCS and bilateral leg or mainly low back pain treated initially with spinal infusions of opioids was then done.

In this study, 42 patients were screened; 26 (62%) patients received spinal stimulation; 16 (38%) received opioids via a spinal infusion pump. Five patients did not receive adequate pain relief with SCS; 3 (7%) of these patients underwent trial spinal infusions and had effective pain relief. There were 4 (10%) patients who underwent a trial of spinal infusion of opioid but did not receive adequate pain relief; these patients were not tested with SCS. Pain severity was rated using a verbal digital pain scale: "On a scale of 0 to 10 where 0 is no pain and 10 is the worst pain you could ever imagine, what is your pain now?" 16/26 patients (62%) had greater than 50% pain relief with SCS. In this study, 2/16 (13%) had greater than 50% pain relief with opioids. Mean follow-up was 2.1 ± 0.3 years. This analysis supports the use of SCS for intractable low back and leg pain.

In this study, 7 (17%) patients suffered complications after implantation of the device; 5 (12%) patients required repositioning of catheter type electrodes and 2 patients required revision of the stimulator generator.

Safety Evaluation

Eleven studies were identified based on the detailed inclusion/exclusion criteria to demonstrate the safety of the PRECISION System. The studies included a total of 1056 patients that were trialed with SCS systems and 880 patients that received implants. The table below depicts the number of patients, the number of events, and the percentage of occurrences of each event compared to the total number of patients. It should be noted that citations cover both Stimulator and RF systems. The clinical experience reported in the literature on RF systems is relevant to determining the safety of totally implantable Stimulator systems.

Table 1: Summary of Risks Identified in the Retrospective Clinical Studies

Risks	# Patients With Adverse Event	Intent-to- Treat Basis N = 1056	Implanted Patient Basis N = 880
Lead Migration	175	16.6%	19.9%
Infection	39	3.7%	4.4%
Epidural Hemorrhage	0	0%	0%
Seroma	0	0%	0%
Hematoma	1	0.1%	0.1%

Paralysis	0	0%	0%
CSF Leak	5	0.5%	0.6%
Over/Under Stimulation, Ineffective Pain Control	46	4.4%	5.2%
Intermittent Stimula- tion	0	0%	0%
Pain Over Implant	16	1.5%	1.8%
Allergic Reaction	6	0.6%	0.7%
Skin Erosion	0	0%	0%
Lead Breakage	35	3.3%	4.0%
Hardware Malfunction	22	2.1%	2.5%
Loose Connection	0	0%	0%
Battery Failure	2	0.2%	0.2%
Other	45	4.3%	5.1%

Clinical Experience-Safety

Clinical data has been collected during a clinical study of the PRECISION System. As of January 15, 2004, 35 subjects were enrolled in the study at multiple sites and 26 subjects had a successful trial stimulation period and were implanted with the PRECISION System. The follow-up period for the 26 implanted patients ranged from two weeks to six months. The following major adverse events were reported.

Table 2: Clinical Experience Safety

Туре	Number of Patients	Resolution
Lead Migration	1	Lead repositioning and subsequent replacement
Output malfunction	1	Device replaced
Infection	1	Infection treated
Pain	1	Lead explanted

Other minor adverse events reported by at least one patient included: receiver malfunction, skin irritation, unpleasant stimulation, CSF leak, infection at implant site, lead migration, and OR cable malfunction. Two of the subjects reported multiple events.

References

- Burchiel, K.J., V.C. Anderson, F.D. Brown, R.G. Fessler, W.A. Friedman, S. Pelofsky, R.L. Weiner, J. Oakley, and D. Shatin. "Prospective, Multicenter Study of Spinal Cord Stimulation for Relief of Chronic Back and Extremity Pain." Spine, 21:2786-2793, 1996.
- Hassenbusch, S.J., M. Stanton-Hicks, E.C. Covington. "Spinal cord stimulation verses spinal infusion for low back and leg pain". Acta Neurochirgica, 64:109-115, 1995.
- Kemler, M.A., G.A.M. Barendse, M. Van Kleef, H.C.W. De Vet, C.P.M. Rijks, C.A. Furnee and F.A.J.M. Van den Wilderberg. "Spinal Cord Stimulation in Patients with Chronic Reflex Sympathetic Dystrophy." New England J of Medicine, 343: 618-24, 2000.
- Kim S. H., R.R. Tasker, and M.Y. Oh. "Spinal Cord Stimulation for Nonspecific Limb Pain versus Neuropathic Pain and Spontaneous versus Evoked Pain." Neurosurgery, 48(5): 1056-1064, 2001.
- Kumar, K., C. Toth, R. Nath, and P. Lang. "Epidural Spinal Cord Stimulation for Treatment of Chronic Pain-Some Predictors of Success. A 15 year experience." Surg Neurol, 50: 110-120, 1998.
- Lang, P. "The Treatment of Chronic Pain by Epidural Spinal Cord Stimulation." AXON, 18(4): 71-73, 1997.
- Ohnmeiss, D., R. Rashbaum, M. Bogdanffy. Prospective Outcome Evaluation of Spinal Cord Stimulation in Patients With Intractable Leg Pain. Spine, 21:1344-1351, 1996.

- Rainov, N.G., V. Heidecke, and W. Burkert. "Short Test-Period Spinal Cord Stimulation for Failed Back Surgery Syndrome." Minim Invasive Neurosurg, 39(2):41-44, 1996.
- Segal, R., B. Stacey, T. Rudy, S. Basser, J. Markham. "Spinal Cord Stimulation Revisited." Neurological Research, 20:391-396, 1998.
- Spieglemann, R. and W.A. Friedman. "Spinal Cord Stimulation: A Contemporary Series." Neurosurg 28:65-71, 1991.
- Villavicencio, A.T., J.C. Leveque, L. Rubin, K. Bulsara, and J.P. Gorecki. "Laminectomy versus percutaneous electrode placement for spinal cord stimulation." Neurosurgery, 46:399-406, 2000.

Contraindications

Patients contraindicated for permanent SCS therapy are those who:

- are unable to operate the SCS system
- have failed trial stimulation by failing to receive effective pain relief
- are poor surgical risks
- are pregnant

Warnings

Heat Due to Charging. Do not charge while sleeping. This may result in a burn. While charging, the Charger may become warm. It should be handled with care. Failure to use the Charger with either the Charging Belt or an adhesive patch, as shown, may result in a burn. If you experience pain or discomfort, cease charging and contact Advanced Bionics.

Magnetic Resonance Imaging (MRI). You should **not** be exposed to Magnetic Resonance Imaging (MRI). Exposure to this diagnostic technology may result in dislodgement of your Stimulator or lead(s), heating of the Stimulator, severe damage to the Stimulator electronics and/or increased voltage through the leads or Stimulator which can cause an uncomfortable or "jolting" sensation.

Pediatric Use. The safety and effectiveness of spinal cord stimulation has not been established for pediatric use.

Diathermy. As an SCS patient, you should not have any form of diathermy either as treatment for a medical condition or as part of a surgical procedure. The high energy and heat generated by diathermy can be transferred through your stimulator system, causing tissue damage at the lead site and, possibly, severe injury or death. The Stimulator, whether it is turned on or off, may be damaged.

Cardiac Pacemakers. Spinal cord stimulators may interfere with the operation of implanted sensing stimulators, such as pacemakers and implantable cardiac defibrillators (ICDs). Be sure your physicians are aware of your spinal cord stimulator before going forward with other implantable device therapies so that medical decisions can be made and appropriate safety measures taken.

Stimulator Damage. Burns may result if the pulse generator case is ruptured or pierced and patient tissue is exposed to battery chemicals. Do not implant the device if the case is damaged.

Posture. Changes in posture or abrupt movements may cause decreases, or uncomfortable or painful increases in the perceived stimulation level. Keep the Remote Control with you at all times, and turn the stimulation down or off before making posture changes. If unpleasant sensations occur, the stimulation should be turned off immediately.

Electromagnetic Interference. Strong electromagnetic fields can potentially turn the stimulator off, or cause uncomfortable or jolting stimulation. Avoid or exercise care around:

• Theft detectors or security screeners, such as those used at entrances/exits of department stores, libraries, and other public establishments, and/or airport security screening devices. It is recommended that you request assistance to

bypass the device. If you must proceed through the device, turn off the stimulator and proceed with caution, and move through the center of the screener as quickly as possible.

- Power lines or power generators
- Electric steel furnaces and arc welders
- Large magnetized stereo speakers

As you approach these devices you may become aware of changing stimulation levels. In rare instances, you could experience an increase in stimulation level to the point that the sensation is uncomfortably strong or possibly "jolting." If this happens, turn off the Stimulator. If the Stimulator suddenly turns off by itself, first move away from the area. Next, check the Stimulator status with the Remote Control by pressing the Stimulation On/Off button and observing the screen. The Stimulator may need to be recharged before stimulation can be restarted. (See "Charging the Stimulator" on page 55 for additional information.)

Always be aware of your surroundings, particularly near theft detectors/security screeners. Ask for assistance to go around these devices if you feel at all uncomfortable.

Precautions

Physician training is required.

Medical Devices/Therapies. The following medical therapies or procedures may turn stimulation off or may cause permanent damage to the Stimulator, particularly if used in close proximity to the device:

- lithotripsy high-output sound or shock waves often used to treat gall stones and kidney stones
- electrocautery the use of a heated electric probe to stop bleeding during surgery
- external defibrillation the use of electrically charged paddles to restart the heart in an emergency
- radiation therapy ionizing energy commonly used to treat cancer
- ultrasonic scanning very high frequency sound waves used to produce images of internal organs or tissue for diagnostic purposes

high-output ultrasound — high frequency sound waves which may be applied as
physical therapy to treat certain bone/muscle injuries, or for muscle stimulation, or to
improve blood flow

Before having procedures, medical therapies, or diagnostics, have your healthcare professional call our Customer Service department at (866) 360-4747 for proper instructions.

Automobiles and Other Equipment. Do not operate an automobile, other motorized vehicle, or any potentially dangerous machinery/equipment with therapeutic stimulation switched on. Turn off stimulation first. Sudden stimulation changes, if they occur, may distract you from attentive operation of the vehicle or equipment.

Post Operative. During the two weeks following surgery, it is important to use extreme care so that appropriate healing will secure the implanted components and close the surgical incisions:

- Do not lift objects of more than five pounds.
- Do not engage in rigorous physical activity such as twisting, bending, or climbing.
- If new leads were implanted, do not raise your arms above your head.

Temporarily, there may be some pain in the area of the implant as the incisions heal. If discomfort continues beyond two weeks, contact your physician.

If you notice excessive redness around the wound areas during this time, contact your physician to check for infection and administer proper treatment. In rare cases, adverse tissue reaction to implanted materials can occur during this period.

Be sure to consult your physician before making lifestyle changes due to decreases in pain.

Stimulator Location. Never attempt to change the orientation or "flip" the Stimulator. Do not "finger" or play with the Stimulator. If the Stimulator flips over in your body it cannot be charged. If you know that the device has turned, or if stimulation cannot be turned on after charging, contact your physician to arrange an evaluation of the system.

In some cases, the skin over your Stimulator may become very thin over time. If this occurs, contact your physician.

Lead Location. In some instances a lead can move from its original location, and stimulation at the intended pain site can be lost. If this occurs, consult your physician who may able to restore stimulation by reprogramming the Stimulator in the clinic or repositioning the lead during another operation.

Device Failure. Stimulators can fail at any time due to random component failure, loss of battery functionality, or lead breakage. If the device stops working even after complete charging (up to four hours), turn off stimulation and contact your physician so that the system can be evaluated.

Storage, Handling and Transport. Do not expose the Remote Control or Charging System components to excessively hot or cold conditions. Do not leave the devices in your car or outdoors for extended periods of time. The sensitive electronics can be damaged by temperature extremes, particularly high heat. For proper operation, do not use the Charger if the ambient temperature is above 35°C (95°F).

If the Remote Control or the Charging System is to be stored for a period of time, be careful that the storage temperature does not exceed -20–60°C (-4–140°F).

Handle the system components and accessories with care. Do not drop them or submerge them in water. Avoid all sources of water that can come into contact with the devices. Although reliability testing has been performed to ensure quality manufacturing and performance, dropping the devices on hard surfaces or in water, or other rough handling, can permanently damage the components. (See "Limited Warranty" on page 73.)

Component Disposal. Do not dispose of the Remote Control or Charger in fire. The battery in these devices can explode in fire. Dispose of used batteries in accordance with local regulations. The Stimulator should be explanted in the case of cremation, and returned to Advanced Bionics. External devices to be disposed of per local regulatory requirements. Please contact your healthcare professional for information.

Remote Control, Charging System Cleaning. The components can be cleaned using alcohol or a mild detergent applied with a cloth or tissue. Residue from soapy detergents should be removed with a damp cloth. Do not use abrasive cleansers for cleaning.

Cell Phones. While we do not anticipate any interference with cell phones, the full effects of interaction with cell phones are unknown at this time. If there is a concern or a problem is encountered, the physician should be contacted.

Adverse Effects

Potential risks are involved with any surgery. The possible risks of implanting a pulse generator as part of a system to deliver spinal cord stimulation include:

- The lead(s) which deliver stimulation may move from their original implanted location, resulting in undesirable changes in stimulation and subsequent reduction in pain relief.
- System failure, which can occur at any time due to random failure(s) of the components or the battery. These events, which may include battery leakage, device failure, lead breakage, hardware malfunctions, loose connections, electrical shorts or open circuits and lead insulation breaches, can result in ineffective pain control.
- Your body may react negatively to the materials used to manufacture the stimulator or the leads. You may notice redness, warmth or swelling of the implant area.
- The skin over your Stimulator may become thin and increasingly tender over time. A seroma may be formed.
- The most common surgical procedural risks are temporary pain at the implant site and infection. However, since the leads are placed in the fluid surrounding your spinal cord, there is a small risk that spinal fluid may leak from the lead insertion site following

surgery. Very rarely, you may develop an internal blood clot (hematoma) or blister (seroma); or you may experience brain hemorrhage or paralysis. Your spinal cord may become compressed.

- External sources of electromagnetic interference may cause the device to malfunction and affect stimulation.
- MRI. Exposure to magnetic resonance imaging (MRI) can result in noticeable heat near the Stimulator or the leads; may distort or destroy the image needed for diagnosis; and may produce enough electromagnetic interference (EMI) to erase the Stimulator programming, destroy the leads, or cause the leads to move from their intended location.
- Undesirable stimulation may occur over time due to cellular changes in tissue around the electrodes, changes in electrode position, loose electrical connections and/or lead failure.
- You may experience painful electrical stimulation of your chest wall as a result of stimulation of certain nerve roots several weeks after surgery.
- Over time, your Stimulator may move from its original position.

- You may experience weakness, clumsiness, numbness or pain below the level of implantation.
- You may experience persistent pain at the Stimulator or lead site.

In any event, you should contact your physician and inform him/her.

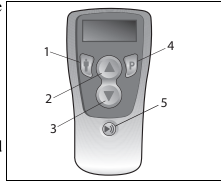
4

The Remote Control

Each button on the Remote Control activates one or more of the stimulation functions which are described in detail on the following pages of this section.

Buttons and Basic Operation

To use most Remote Control functions, you will simply press a button as you would on a TV remote control. Other functions require a "long press," which is described in "Selecting Options" on page 47.



- 1. Area Button
- 4. Program Button
- $2.\ Up/Activate$
- 5. Stimulation On/Off
- 3. Down/Save

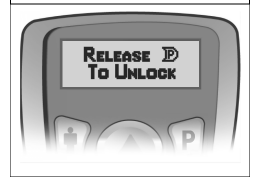
Basic Operation

When it is not being used, the Remote Control sets itself in a "sleep" or idle mode and the display screen is blank. During this sleep mode when you press any button *except* the **>)** button (stimulation on/off), the Remote Control will "wake up" and display the screen shown on the right.

To unlock the Remote Control, press and hold **P**, until "Release **P** To Unlock" appears. The Remote Control will immediately look for your Stimulator and then connect with it, allowing you to make adjustments to your stimulation.

When you are finished using it (no buttons are being pressed), the Remote Control will return to the sleep mode within a minute or so.



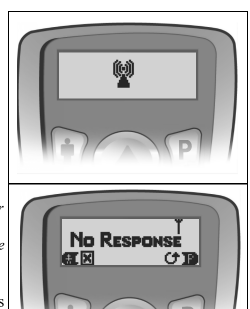


Communicating with the Stimulator

Good communication between your Stimulator and the Remote Control is very important. For that reason, you will sometimes see the following icon on your screen. This indicates that the Remote Control is checking for the Stimulator.

Note: If there is a problem communicating with the Stimulator, the message "No Response" will appear on the Remote Control screen. Press to retry, or press to cancel. See "Help" on page 65 for more information.

The Remote Control unit is your direct link to choices available for tailoring spinal cord stimulation to suit your comfort and convenience requirements. *Keep the Remote Control with you at all times*.

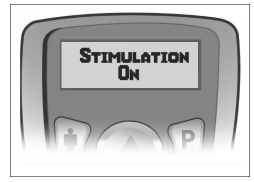


Stimulation On and Off

The Remote Control uses a "dedicated" stimulation on/off switch. You may press •) at any time—even during the Remote Control's sleep or idle mode—to turn stimulation on or off. The Remote Control will briefly display a message notifying you of the on or off status.

Signal Strength

The Remote Control indicates the signal strength between the unit and the Stimulator in the upper right corner of the display:





- If there are no signal bars, move the Remote Control closer and tilt it. If there are no signal bars, communication may still be achieved, however, it is at a very weak level.
- If there are 1 or 2 signal bars, there is adequate communication between the Remote Control and the Stimulator

- If there are 3 or 4 signal bars, the Remote Control is achieving optimum communication with the Stimulator.
- After missing a telemetry message, the Remote Control will display the "Searching" message. The Remote Control will then start looking for signal strength every second. Move to a better spot and wait for the signal strength to display. To cancel searching press †.

Note: Avoid common sources of interference, such as televisions and computer monitors.

The Level Screen

Whenever stimulation is turned on, or after the Remote Control is awakened from sleep/idle mode, the Remote Control display defaults to the Level screen. When you press the ▲ or ▼ button from this display, you are able to increase or decrease the stimulation strength, or level, of *all* of your pain areas at once (if your Stimulator was programmed to treat more than one area). To learn about adjusting



the stimulation of individual areas, please see "Selecting Areas for Level Control" on page 38.

The Level screen also displays a bar graphic near the top center to indicate the battery charge level of your Stimulator.

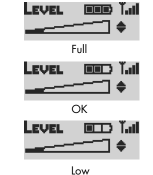
Note: This is the battery level of your Stimulator, not your Remote Control.

The graphic is easy to understand: Three filled-in bars means that the Stimulator has a fully-charged battery. As the battery strength wears down, depending on your stimulation settings and usage, the bars will "empty" accordingly. For complete information on maintaining your Stimulator's battery for uninterrupted delivery of the therapy prescribed for you, see "Charging the Stimulator" on page 55.

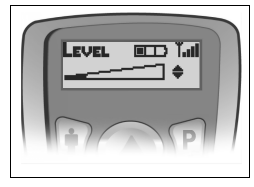


The Remote Control unit is your direct link to choices available for tailoring spinal cord stimulation to suit your comfort and convenience requirements. *Keep the Remote Control with you at all times*

Note: Remember, the Level screen will remain on the display for approximately one minute if you do not press a button to perform an action with the Remote Control. After a minute, the Remote Control will go into idle mode and the display will go blank.



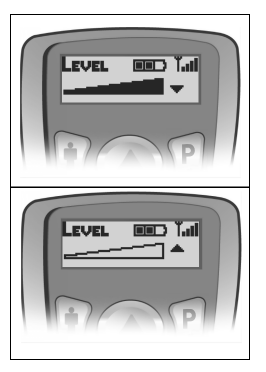
• Multiple area control is available only if your system has been programmed to deliver stimulation to separate areas. If you do not have separate area control (for example, left leg vs. right leg) but feel that separate control might improve your therapy, contact your healthcare professional to determine what is possible.



Stimulation Level Control

Occasionally, you may see one of the accompanying messages on the Remote Control screen while changing the stimulation level. These displays, and a "beep" from the Remote Control, alert you that you have reached either a maximum or a minimum level:

The bar replacing the up arrow means that you have reached the maximum allowable level and can only *decrease* the stimulation setting. The bar replacing the down arrow means that you have reached the minimum level and can only *increase* the stimulation setting.



Selecting Areas for Level Control

Your Remote Control may have as many as four therapy areas stored for your use. Each area will have a name or a number (1 through 4) which was assigned during your programming at the clinic. The stimulation level for each of these areas is controlled from an individual "area level" screen. To change the stimulation strength of a single area:

1. From the main Level screen, press the button as many times as necessary to cycle through your programmed area Level screens.



Note: Button presses will move from one area to the next and will eventually return you to the Level screen.

2. When the named or numbered area that you want to adjust is shown on the screen, press ▲ or ▼ to change the stimulation level for that area.

The screen will go blank and the Remote Control will go to "sleep" after one minute of inactivity.

Programs

A stimulation program may provide paresthesia to cover a single pain area, or up to four areas, depending on how the program was set up by your clinician. Your Remote Control can store up to four programs—numbered 1 through 4—for you to select and activate at any time. Often, the programs will have certain differences in the settings to allow you to vary your stimulation in several ways. You may have been encouraged to try using specific programs for different circumstances, or different body positions, or different times in your daily routine. Programs and their flexibility give you and your healthcare professional a way to continually "fine tune" your therapy.

Selecting and Activating Programs

1. Press the **P** button from the Level screen to go to the Program screen.

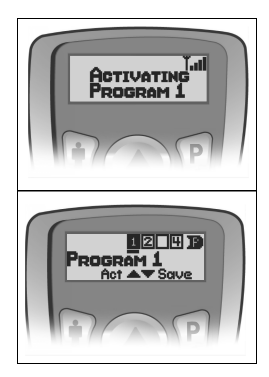
From the Program screen, each additional **P** button press selects the next program number. Press † to select the previous program number. The black highlight box shows where you are in the cycle. Pressing **P** from program 4 returns you to the Level screen. Pressing † from program 1 also returns you to the Level screen.



2. Press P or to move until the program you want is highlighted, then press ▲ to activate the program.

Note: You might not have four programs saved in your Remote Control. Empty program slots will have a blank box with no program number. If you try to activate an empty program, nothing will happen.

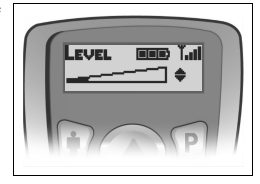
 An underline (_) beneath a program number means that program was the most recently activated or saved (see below).



Once you have selected and activated a program, the Remote Control will return to the Level screen. Use the ▲ or ▼ to adjust the stimulation level of the program, if you wish.

Saving Program Changes

If you do make a stimulation level adjustment and decide that you prefer it to the original setting, you can save the new level (or any other change) and make it a permanent part of the program:



- 1. After making a change to a program, press **P** from the Level screen to return to the Program screen.
- 2. From the Program screen, press **P** as many times as necessary to select the active program; that is, the program that is currently running and the one you have just changed.
- 3. With the active program selected, press ▼ to save the change in the Remote Control's memory.



The Remote Control will first ask you to confirm that you want to "overwrite" the program; press the appropriate button.

If you confirm the change by pressing \mathbf{P} , the program will be updated and saved. If you decide not to make a permanent change, pressing $\mathbf{\uparrow}$ will return you to the Program screen.

You may also save a program change as a completely new program if an empty program slot [] is available:

Simply select the empty slot using **P** and press **▼**. After saving the new program, the Remote Control will return to the Level screen.



Options

Under some circumstances, and depending on your treatment prescription, your healthcare provider may have given you therapy control beyond selectable programs by making special *options* available to you. Program options make it possible for you to change certain preset stimulation settings, and/or return changed programs to their original clinic settings. The ability to go back to original settings allows you to change your mind... then change your mind again... about how satisfied you are with your stimulation paresthesia.

If you have been told about program options and instructed in how to use them, you may realize that you probably will not "exercise" your option settings very often. However, if your Remote Control has been set up to access options, please follow the steps detailed in "Selecting Options" on page 47 to make adjustments.

There are three *possible* options. One of these, Restore, is not a stimulation setting but is similar to an "undo" feature. It returns a changed program back to its original settings. The other two options are stimulation settings that can affect the overall feeling of the stimulation you receive:

- *Pulse Width*, or how long each stimulation pulse lasts, and
- *Rate*, or how many times-per-second your Stimulator sends a stimulation pulse.



Note: Restore is available to all patients. However, Pulse Width and/or Rate may not be available to you. Their availability would have been discussed with you at your first programming session.

Selecting Options

To access the options that may be available through your Remote Control, you will need to use a "long" button press. This simply means that you will *press and hold* a button until a particular option screen is displayed. The action you take next depends on which option you want to adjust. Follow the appropriate instructions below:

To Adjust Stimulation Pulse Width:

- 1. From the Level screen, press and hold

 until you see the Width Area 1 screen. If this is the area you want to adjust, press either

 or ▼ to increase or decrease the Pulse Width.
- 2. **To move to another area**, press briefly (a "normal" press) to cycle through your programmed areas until you find the Width Area screen (2, 3 or 4) that you want.
- 3. Press either ▲ or ▼ to increase or decrease the Pulse Width



Note: The Remote Control will beep to notify you if you reach a preset limit while increasing or decreasing the Pulse Width.

To return to the Level screen from any Pulse Width screen, press and hold † until Rate Area 1 is displayed, then press and hold † again to see the Level screen. If you do not have access to the Rate option, the Level screen will appear right away.

To Adjust Stimulation Rate

1. From the Level screen make a long • button press to Pulse Width Area 1.

Note: If you do not have access to the Pulse Width option, the long press will take you directly to the Rate Area 1 screen.



- 3. To move to another area, press † briefly (a "normal" press) to cycle through your programmed areas until you find the Rate Area screen (2, 3 or 4) that you want.
- 4. Press either ▲ or ▼ to increase or decrease the Rate.
- **Note:** You may also find Rate Area 1 by making a long button press from any Pulse Width screen.
 - The Remote Control will beep to notify you if you reach a preset limit while increasing or decreasing the Rate.

To return to the Level screen from any Rate screen, press and hold • .

To Restore a Clinician Program

Over time, you may have made changes to one or more of the programs originally saved to your Remote Control. The Restore option allows you to return to those original stimulation settings if ever you become dissatisfied with a changed program.

1. From the Level screen, press and hold **P** to reach the Restore screen. Program 1 will be highlighted.

Note: The highlighted program is the current selection; an underline (_) indicates the last program restored; a [] indicates an empty program slot.

2. If necessary, press P to cycle through the program numbers to select the program you want to restore. When the desired program is highlighted, press ▲. The Remote Control will briefly flash a message confirming the restoration.





Understanding Battery Messages

About the Remote Control Batteries

As an SCS patient, it is essential that you appreciate the importance of battery power! The replaceable batteries in your Remote Control and the rechargeable battery in your stimulator work together to provide you with consistent, dependable paresthesia. Always pay close attention to the battery status messages described in this section.



When the Remote Control batteries are at a low power level, the battery message shown here will be displayed. It is a good idea to replace the batteries at this point. *If you continue to use the Remote Control without replacing the batteries, you will eventually see the message "Replace Remote Battery."* Do not ignore the message to replace the batteries.

When you press the **P** button from this screen, the Remote Control will make a check of your Stimulator's battery status also.

If the Stimulator battery is full or OK, the Remote Control will return to the Level screen. To learn about the messages you will see when the Stimulator's battery is less than full, see page 53.

If you do not respond to the Remote Battery Low message in a timely manner, the batteries will eventually drain to the point of not having enough power to manage your stimulator.

Next, you would see the more urgent message shown opposite. You must respond immediately! When you press **P**, the Remote Control will make another check of your Stimulator battery and may display one of the messages shown on the following page.

Note: Make a habit of replacing the Remote Control batteries when you first see the Remote Battery Low message. See "Battery Replacement" on page 54.



About Your Stimulator Battery

When the Remote Control checks the condition of your Stimulator battery, you may see one or more of the following messages depending on the battery's charge level:

- If the Stimulator battery is low, the Remote Control will display the message shown at right.
 Press P to return to the Level screen.
- If the Stimulator battery is *very* low, the Remote Control will display this notice to recharge the stimulator battery immediately. Stimulation has most likely been turned off automatically.

Note: The Remote Control will continue to display this message until the stimulator is recharged.



Battery Replacement

- 1. On the rear of the Remote Control, push in slightly and slide down the battery compartment cover.
- Remove the old batteries.
- 3. Place the three new AAA batteries in the slots, matching the positive (+) and negative (-) markings in the compartment.
- 4. Align the battery compartment cover on the case

 and slide the cover into position until it snaps closed.



The Remote Control will connect to the Stimulator in approximately 30 seconds.

Note: It is recommended that you remove the batteries if the Remote Control is not going to be used for an extended period of time.

5

Charging the Stimulator

Your Precision Stimulator uses a rechargeable battery to provide stimulation. The Precision System's programming software gives your healthcare provider a conservative recommendation for how often to charge. This estimate is based on 24 hours per day/7 days a week at the maximum energy output level. While you may want to follow these recommendations, you can also develop a charge routine that fits best into your own lifestyle. Developing a recharge schedule involves finding the right balance between four things:

- How much power you need in order to have effective therapy.
- How often you want to recharge.
- How long you want to recharge.
- How you would like to manage your personal schedule.

In most cases, the Stimulator will need to be recharged at intervals ranging from once a week to once a month. The Remote Control provides an easy-view Stimulator battery charge status graphic on the Level screen as well as messages to inform you of the battery's condition. These messages are explained beginning on page 53.

Following the general recommendations made by your healthcare provider, you have the freedom to establish the charging routine that suits you best. Keep in mind that if you do not charge your Stimulator before it reaches the very low "hibernation" condition, stimulation will eventually stop until you charge again. Developing a charging routine you are comfortable with will help to prevent you from losing stimulation due to a low battery.

Getting Started

WARNINGS:

- Do not charge while sleeping. This may result in a burn.
- While charging, the Charger may become warm. It should be handled with care.
- Failure to use the Charger with either the Charging Belt or an adhesive patch, as shown, may result in a burn. If you experience pain or discomfort, cease charging and contact Advanced Bionics.

The Charging System for your Stimulator consists of the Charger unit, a Base Station, and a Power Supply. The Base Station is designed to remain connected to a power outlet at all times. When it is not being used, keep the Charger on the Base Station so that it is always ready to deliver a charge.

- 1. Find a convenient place with a flat, clear surface to keep the Base Station plugged in.
- 2. Plug the Power Supply into a standard AC wall outlet, and then plug the other end into the Base Station connector.

3. Place the Charger in the Base Station.

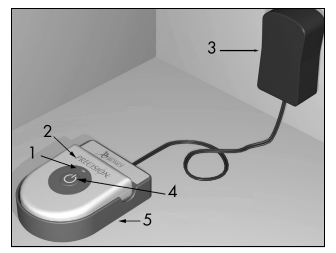
When charging, you must use the Charger with either the Charging Belt or an adhesive patch to hold the Charger over the Stimulator. Before using the Charging Belt, you may cut it to a more convenient size, but be careful not to cut too much. The adhesive patches are made of non-reactive material suitable for most sensitive skin types.

Note: Advanced Bionics recommends the use of the Charging Belt for charging.

The Charger is completely ready and able to fully charge your Stimulator when the indicator light is green. If the light is amber-colored, the Charger can only partially charge the Stimulator. It may be used, but it may not be able to return your Stimulator to a full charge (and you may need to charge sooner than you normally would).

Indicator Light Status:

- Green ready for full charge
- Amber partial charge
- Off not ready for charging



- 1. Indicator Light
- 2. Charger
- 3. Power Supply

- 4. Power Button
- 5. Base Station

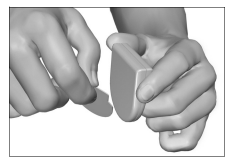
Charging Your Stimulator

- 1. When the indicator light is green, remove the Charger from the Base Station. (*The indicator will go off, regardless of the ready status of the Charger*.)
- 2. Using the Adhesive Patch:
 - Apply the adhesive patch to the backside of the Charger by peeling the clear liner from the patch and applying the white side with the blue stripe to the rear of the Charger, as shown. (See diagram below.)
 - Remove the skin side beige liner from the adhesive (only good for one fixation.) (See diagram below.)

OR

Using the Charging Belt:

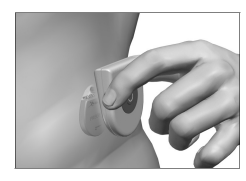
• Place the Charger into the pocket on the Charging Belt with the power button facing out. (See diagram below.)





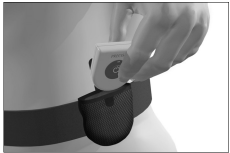
3. Press the power button. The indicator light will come on again, and the Charger will begin beeping steadily to signal that it is searching for the Stimulator.

4. Locate the Charger over the Stimulator. When the Charger is aligned with the Stimulator, the beeping will stop.



5. Secure the Charger over the Stimulator either by pressing the adhesive to the skin over the Stimulator, or by securing the Charging Belt.

Note: If you accidentally locate the patch in the wrong place, or if the Charging Belt moves out of alignment, the Charger will start beeping again. Use a new adhesive patch or readjust the belt to place the Charger back into position.



6. When the Charger emits a distinct double beep, the Stimulator is charged. Switch off the Charger, remove the Charging Belt or adhesive patch, and return the Charger to the Base Station.

Do not confuse the end of charge signal (a distinct double beep) with the steady, continuous misalignment signal.



- **Note:** Depending on your program parameters, you may expect daily recharging times from as low as 10 minutes up to four hours, or weekly recharging times from as low as one hour up to four hours.
 - The end of a charge signal is a distinct double beep, and the alignment indicator is a steady continuous signal.

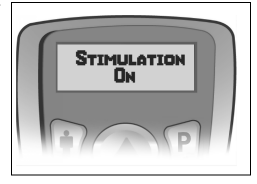
The rechargeable Stimulator battery should provide you with at least five years of service. Over time and with repeated charging, the battery in your Stimulator will lose the ability to recover its full capacity. As a result, you may need to recharge your Stimulator for longer periods and/or more often. Your Stimulator will need replacement when stimulation can no longer be maintained with routine daily charging.

6 Help

Stimulation

No Stimulation

- 1. Toggle the N Remote Control stimulation on/off button to make sure that stimulation is ON. If the Remote Control receives confirmation from the Stimulator, it will flash "Stimulation On."
- 2. Turn up the level of stimulation from the main screen or area screens.
- 3. Charge the Stimulator. When the charge is complete, try turning the stimulation on.



4. Call our Customer Service Department at (866) 360-4747 if the above steps do not solve the problem.

Stimulation Increases or Decreases on Its Own

- 1. Stimulation can change depending on body position (lying down, standing or bending).
- 2. Always keep the Remote Control with you, so that you can adjust your stimulation levels as needed.

Stimulation Shuts Off

- 1. When the Stimulator battery needs to be recharged, it will stop stimulating. Check the battery status with the Remote Control and recharge if necessary, then turn stimulation back on. If the Stimulator regularly stops stimulating before you charge, you can charge more often.
- 2. Although unlikely, anti-theft screeners can turn stimulation off. If you cannot turn the stimulator back on with your Remote Control, you may need to charge the Stimulator.

3. Large magnetized speakers or large power lines that emit interference may also turn off stimulation. If you cannot turn the stimulator back on with your Remote Control, you may need to charge the Stimulator.

Remote Control Display Messages

"Remote Battery Low" on the Display

The batteries in your Remote Control should be replaced soon. All of the Remote Control functionality is still available (refer to "Battery Replacement" on page 54).

"Replace Remote Battery" on the Display

The batteries in your Remote Control need to be replaced with three fresh AAA batteries in order to have full functionality of the Remote Control (refer to "Battery Replacement" on page 54).



Searching

"Searching" on the Display

In the event of communication interference, the Remote Control will automatically begin "searching" for the Stimulator. Try to reposition the Remote Control closer to the Stimulator to help it locate the Stimulator.

"No Response" on the Display

When the Remote Control displays "No Response," there is a communication problem between the Remote Control and the Stimulator. Press the button to activate automatic searching for the Stimulator. Then reposition the Remote Control until it beeps. This indicates that communication has been established



Once the Remote Control connects with your Stimulator, you will be returned to the display you were using before the problem began. If the problem persists, replace the Trial Stimulator battery and check to see if the problem is solved.

Occasionally, telemetry problems occur because orientation or interference prevent the Remote Control from finding the Stimulator. Move the Remote Control closer to your Stimulator and then press the **P** button.

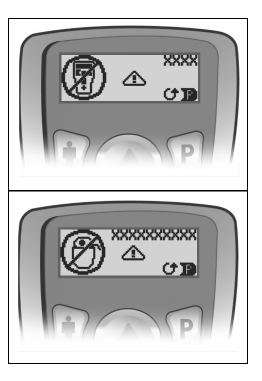
Call our Customer Service Department at (866) 360-4747 if the problem continues.

"Error Code" on the Display

If the Remote Control displays the error screen shown at right, try to make a note of the numbers (the error code) on the top line. Then press **P**, wait a few seconds, move the Remote Control closer and try the action again.

Since most of the error codes you might encounter are related to communication, always try to resolve the problem by moving the Remote Control closer to the Stimulator. If this does not work, please contact our Customer Service Department.

Note: The Remote Control will go to the Level screen when \mathbf{P} is pressed, or to idle mode if \mathbf{P} is not pressed within one minute.



Accessories

Washing the Charging Belt

Wash the Charging Belt with mild soap and warm water.

Contacting Advanced Bionics

If you have any other questions, or need to contact Advanced Bionics for any reason, you may do so in any of the following ways:

Customer Service Phone: (866) 360-4747

• Customer Service Fax: (661) 362-1503

Address: Advanced Bionics Corporation

Pain Management Division

Mann Biomedical Park 25129 Rye Canyon Loop

Valencia, CA 91355

7

Limited Warranty

Implanted Pulse Generator

Advanced Bionics[®] Corporation (referred to as Bionics) warrants to the patient who receives a Precision[®] Implantable Pulse Generator (referred to as the IPG) that the IPG will be free from defects in workmanship and materials for a period of (5) five years from the date of surgical implant of the IPG. This warranty applies only to the patient who has the IPG implanted and no other person or entity. This warranty does not apply to the leads, extensions, or surgical accessories used with the IPG.

If the IPG fails to function within normal ranges within (5) five years after the date it is implanted, Bionics will replace the IPG with a functionally equivalent IPG made by Bionics. No other relief whatsoever is available under this limited warranty. The limited warranty for a replacement IPG will last only for five years from the date of surgical implant

of the original IPG. Claims under this limited warranty are subject to the following additional conditions and limitations:

- 1. The product registration card must be completed and returned to Bionics within 30 days of surgery.
- 2. The IPG must be purchased after January 1, 2005 and implanted before the "use before" date.
- 3. Failure of the IPG must be confirmed by Bionics.
- 4. The IPG must be returned to Bionics (or Bionics' authorized agent) within 30 days after it fails to function within normal ranges. That IPG will be Bionics' property.
- 5. This limited warranty does not include failures to function within normal ranges caused by:
 - (a) fire, floods, lightning, natural disasters, water damage and other calamities commonly defined as "Acts of God";
 - (b) accident, misuse, abuse, negligence, or the customer's failure to operate the IPG in accordance with manufacturer's instructions;

- (c) unauthorized attempts to repair, maintain, or modify the IPG by the patient or any unauthorized third party; or
- (d) attaching equipment to the IPG that is not supplied or expressly authorized by Bionics.

THIS LIMITED WARRANTY IS THE ONLY WARRANTY THAT APPLIES TO THE IPG, AND BIONICS EXPRESSLY DISCLAIMS ANY OTHER WARRANTY, EXPRESS OR IMPLIED, INCLUDING ANY WARRANTY OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

Under this limited warranty, Bionics will be responsible only for replacement of the IPG with a functionally equivalent IPG made by Bionics and will not be liable for any damages (whether direct, indirect, consequential, or incidental) caused by the IPG, whether the claim is based on warranty, contract, tort or any other theory.

Externals

Advanced Bionics warrants to the patient that the Remote Control device and Charging System (Charger and/or Charger Base Station) are free from defects in workmanship and materials for a period of one (1) year from the date of purchase.

If a Remote Control Device or Charging System component fails to function within normal ranges within one year after the date of purchase, Bionics will replace the device or component with a functionally equivalent device or component made by Bionics. No other relief whatsoever is available under this limited warranty. The limited warranty for a replacement device or component will last only for one year after the date of purchase. Claims under this limited warranty are subject to the following additional conditions and limitations:

- 1. The product registration card must be completed and returned to Bionics within 30 days of purchase.
- 2. Bionics must confirm the device or component failure.
- 3. The device or component must be returned to Bionics (or Bionics' authorized agent) within 30 days after it fails to function within normal ranges. That device or component will be Bionics' property.
- 4. This limited warranty does not include failures to function within normal ranges caused by:
 - (a) fire, floods, lightning, natural disasters, water damage and other calamities commonly defined as "Acts of God";

- (b) accident, misuse, abuse, negligence, or the customer's failure to operate the device or component in accordance with manufacturer's instructions;
- (c) unauthorized attempts to repair, maintain, or modify the device or component by the patient or any unauthorized third party; or
- (d) attaching equipment to the device or component that is not supplied or expressly authorized by Bionics.

THIS LIMITED WARRANTY IS THE ONLY WARRANTY THAT APPLIES TO THE DEVICE OR COMPONENT, AND BIONICS EXPRESSLY DISCLAIMS ANY OTHER WARRANTY, EXPRESS OR IMPLIED, INCLUDING ANY WARRANTY OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

Under this limited warranty, Bionics will be responsible only for replacement of the device or component with a functionally equivalent device or component made by Bionics and will not be liable for any damages (whether direct, indirect, consequential, or incidental) caused by the device or component, whether the claim is based on warranty, contract, tort, or any other theory.

Glossary

- Area Button icon. This figure respresents the area button on the Remote Control.
- **ADHESIVE PATCH.** Non-reactive skin patch designed to temporarily attach the Charger to the skin over the Stimulator site.
- ADVERSE EFFECT. Undesirable result.
- **AMPLITUDE.** The measure-of-strength of delivered stimulation. (See Level.)
- **AREA.** A location on the body such as right leg or left leg where stimulation will occur.
- **CARDIAC PACEMAKER.** A small implantable device used to control the rhythm of the heart.

- **CHARGER.** A portable device used to recharge the battery of the implanted stimulator.
- CHARGER BASE STATION. A holder/ power supply that supports the Charger and keeps it in a ready state for recharging the Stimulator.
- CHARGING SYSTEM. The Charging System consists of a Charger Base Station, Charger, Power Supply, Charging Belt and adhesive patches. The system is used for recharging the implanted stimulator.
- **CONTROL BUTTONS.** Buttons located on the Remote Control; used for adjusting stimulation settings.
- **DIATHERMY.** A therapeutic procedure used to heat body tissue by high-frequency electromagnetic currents.

- **DISPLAY.** The Remote Control screen.
- **ELECTRICAL PULSE GENERATOR.** Also called an implantable pulse generator (IPG or Stimulator); used to send electrical pulses to the spinal cord or other parts of the body.
- **ELECTRICAL STIMULATION.**The energy created by a pulse generator.
- **ELECTROMAGNETIC INTERFERENCE** (EMI). Electromagnetic signals that interfere with a variety of electrical signals including spinal cord stimulation.
- **IDLE MODE.** A time-out period when the Remote Control is not being used. Also known as sleep mode.
- **INCISION.** Small surgical cut or opening in the skin.

- **INDICATOR.** A signal light used on the Trial Stimulator and the IPG Charger.
- **IPG.** Implantable Pulse Generator.
- **LEAD.** A surgical wire that sends electrical stimulation pulses from a pulse generator to the spinal cord.
- **LEAD MIGRATION.** The movement of a lead away from the spinal cord.
- **LEVEL.** Term used on the Remote Control screen to identify the amplitude or strength of stimulation pulses.
- **LONG BUTTON PRESS.** To press and hold a button for about 3 seconds.
- **MRI.** Magnetic Resonance Imaging; the use of a nuclear magnetic resonance spectrometer to produce electronic images of tissues and organs.

- **OPTIONS.** Methods for adjusting stimulation beyond amplitude, or level, using the Remote Control. Your healthcare provider may or may not provide you with all available options.
- Program Button icon. This symbol represents the Program button on the remote control.
- **PARESTHESIA.** Sensation produced by electrical stimulation.
- PATIENT IDENTIFICATION CARD. A wallet size card that lists the patient name, physician name, Stimulator model and serial number.
- **PERMANENT IMPLANT.** A stimulator system, pulse generator and leads, implanted in the body and maintained by a pulse generator battery Charging System.

- **PRECAUTION.** Generally, situations that you should be aware of in order to avoid potentially uncomfortable stimulation sensations and/or damage to your stimulation system.
- **PROGRAM.** Combination of one or more stimulation areas.
- **PULSE WIDTH.** The length of time each stimulation "spark" lasts. An option setting available from the Remote Control.
- **RATE.** The number of times—per-second (speed) at which stimulation pulses are delivered to the spinal cord. An option setting available from the Remote Control.
- **REMOTE CONTROL.** A battery powered hand-held computer used to adjust stimulation.

- **SAVE.** The Remote Control button command used to store a newly created or modified stimulation program.
- **SLEEP MODE.** A time-out period when the Remote Control is not being used. Also known as idle mode.
- SPINAL CORD STIMULATION (SCS). A method of applying electrical pulses to the spinal cord to block/mask pain signals to the brain.
- **STIMULATION.** When used as a therapy for pain, an artificially applied, low-level, pulsating electrical "shock" felt as a tingling or pulsating sensation in the area of pain and perceived enough to reduce the awareness of pain.
- **STIMULATION COVERAGE.** Area on the body where stimulation occurs. (See Area.)

- **STIMULATOR.** Small implantable electrical pulse generator used to control stimulation.
- d or ▶) Stimulation On/Off Button icon.
- **SUBCLAVICULAR.** Under the collarbone.
- **SYSTEM FAILURE.** Inability of spinal cord stimulator system to deliver stimulation therapy.
- **WARNING.** Potential hazards that you must be aware of to avoid serious situations that may cause injury or death.

WIDTH. See Pulse Width.

Index

A	
Adhesive Patch 57, 73	;
amplitude 73, 74, 75	,
area)
В	
battery	ļ
C	
charge indicator 56, 58	3
charge signal60)
control buttons	;
D	
diathermy	3

S

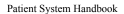
SAVE76

The following is federal government communications regulation information about the Precision[™] Spinal Cord Stimulation System.

This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions: (1) This device may not cause harmful interference, and (2) This device must accept any interference received including interference that may cause undesired operation.

The Precision System components should only be serviced by Advanced Bionics. Do not attempt to open or repair any of the components. Unauthorized opening of or attempts to repair the components will void the warranty.

Changes of modifications to this product not authorized by Advanced Bionics Corporation could void the FCC Certification and negate your authority to operate this product.





CORPORATE HEADQUARTERS

Advanced Bionics[®] Corporation 12740 San Fernando Road, Sylmar, CA 91342 (800) 678-2575 in US and Canada (818) 362-7588, (818) 362-5069 Fax (800) 678-3575 TTY www.advancedbionics.com Email:info@advancedbionics.com

PAIN MANAGEMENT DIVISION

Advanced Bionics[®] Corporation Mann Biomedical Park 25129 Rye Canyon Loop, Valencia, CA 91355 (661) 362-1400, (661) 362-1500 Fax

Part No. 9055520-001 Rev A

©2006 Advanced Bionics Corp. All rights reserved.