

MARGINPROBE®



User Manual

User Manual P/N: PB0501071

This page is intentionally left blank

COPYRIGHT

© 2012 Dune Medical Devices Ltd. All rights reserved worldwide.

This work and all rights herein are owned exclusively by Dune Medical Devices Ltd., its subsidiaries and its other affiliated corporations. No parts of this manual, in whole or in part, may be reproduced, performed, published, displayed, broadcast, translated, or transmitted to any electronic medium or machine readable form, distributed, sold or otherwise used or relied upon without the express prior written permission of Dune Medical Devices Ltd. Reproduction or reverse engineering of copyrighted software is prohibited.

PATENTS

This product is protected by the following US Patents: 7,899,515; 7,904,145; 8,019,411; 8,147,423; 8,195,282; 8,319,502. This product is protected by the following Chinese Patent: ZL200680019026.4. Other patent applications are pending in the USA, China, Japan and Europe, including: EP1890596; EP1919273; EP2118801; EP2129281; EP2304456.

TRADEMARKS



• *Medical Devices* • is a registered trademark of Dune Medical Devices Ltd.

• DUNE is a registered trademark of Dune Medical Devices Ltd.

• MARGINPROBE is a registered trademark of Dune Medical Devices Ltd.

• All product names and any registered and unregistered trademarks mentioned in this manual that refer to goods or services offered by Dune Medical Devices Limited, its subsidiaries and other affiliated companies are owned by Dune Medical Devices Ltd.

• All third-party product names and any registered or unregistered trademarks mentioned in this manual that refer to goods or services offered by parties other than Dune Medical Devices Limited remain the property of their respective owners.

User Manual Part Number:	PB0501071
Revision Level:	А
Revision Date:	January 2013
This User Manual refers to:	MarginProbe [®] System - Model SA0580000

Contact information for equipment manufacturer and technical services:

Dune Medical Devices Inc.

25 Thomson Place, Suite 440 Boston, Massachusetts 02210 USA Tel: +1-508-620-2782 info@dunemedical.com This page is intentionally left blank

TABLE OF CONTENTS

1. Before You Start Conventions Used in this Manual	1-1 1-1
2 Safety and Regulatory Issues	
Intended Lise	2 1 2 1
Warnings Processions and Contraindications	
Potential Adverse Effects of the Device on Health	۲-۲-۲۲۰
Clinical Data	د-∠
Electrical and Mechanical Salety Precautions.	
Compliance with international Regulatory Standards	
Device Regulatory Labels	
Probe identification Label	
EtO Exposure Indicator	
3. Console Placement	
Facility Requirements	3-1
Electrical Requirements	3-1
Space Requirements	3-1
Environmental Requirements – MARGINPROBE	3-3
Storage Requirements – MARGINPROBE [®] Console	
Storage Requirements – MARGINPROBE® Probe	
Console Placement	
4. System Description	4-1
Introduction	4-1
Console	4-2
Control Panel	4-3
Summary of control panel buttons	4-3
User Interface	4-4
Probe Components	
Coupling and Measurement	
Probe Control Button	4-6
Probe LEDs	4-7
Probe Connector	4-7
5. Operating Instructions	
Introduction	
Preparing the System for Operation	
Connect the Probe outside the Sterile Field.	
Operating the System	5-4
Probe Identification	5-5
Grouping Measurements According to Margins	5-6
Performing Measurements	
Interpreting Measurement Results	5_10
Screen Scrolling	5-10 ج 14
Drohe Lleage Limitations	۲۱۱-ن-ن ۲۹۵ ه
FIDDE USAYE LIITIILAIIUTS	

Clearing the Screen and Removing Popup Messages	5-12
Standby Mode	5-12
System Shut-Down	5-13
6. Maintenance	6-1
Introduction	6-1
Exterior Cleaning	6-1
Interior Inspection	6-1
Service Information	6-1
Storage	6-2
7. Troubleshooting	7-1
Introduction	7-1
Troubleshooting Guides	7-1
8. System Specifications	8-1
System Performance	8-1
Environmental Requirements	8-1
Physical Specifications	8-1
Electrical Specifications	8-1

LIST OF FIGURES

Figure 2-1: Pivotal Study Design	2-5
Figure 2-2: Illustration of CSR Primary Endpoint	2-7
Figure 2-3: CSR and Clinical Relevance	2-8
Figure 2-4: Pivotal Study Patient Flow Chart - SOC Arm	2-22
Figure 2-5: Pivotal Study Patient Flow Chart - SOC+Device Arm	2-23
Figure 2-6: Algorithm Development Process	2-26
Figure 2-7: Study III - ROC curves of 3 different datasets	2-28
Figure 2-8: Study V - Sampling Process	2-30
Figure 2-9: Study V - Final Pathologies from Patients Treated at Study Sites 1 and 2 (Selective Re- excision)	2-31
Figure 2-10: Study V - Final Pathologies from Patients Treated at Study Site 3 (Total Re-excision)	2-31
Figure 2-11: MAST Study - Final Pathologies - SOC Arm	2-32
Figure 2-12: MAST Study - Final Pathologies - SOC+Device Arm	2-33
Figure 3-1: System Dimensions	3-2
Figure 3-2: Electrical Connection	3-4
Figure 4-1: MARGINPROBE [®] System	4-1
Figure 4-2: System Description: Console	4-2
Figure 4-3: System Description: Control Panel	4-3
Figure 4-4: Graphic User Interface	4-4
Figure 4-5: System Description: Probe	4-5
Figure 4-6: Probe Tip	4-6
Figure 4-7: Control Button and Probe LEDs	4-6
Figure 4-8: Probe Connector	4-7
Figure 5-1: Connect Probe Popup Window	5-2
Figure 5-2: Standby Mode Message	5-2
Figure 5-3: Probe Connection	5-3
Figure 5-4: Calibrating Message	5-4
Figure 5-5: New Probe Screen	5-5
Figure 5-6: choosing the Margin to be measured	5-6
Figure 5-7: Confirmed Chosen Margin	5-7
Figure 5-8: Probe Tip Application to the Specimen	5-9
Figure 5-9: Measurement Results	5-11
Figure 5-10: Screen Scrolling	5-11
Figure 5-11: Shutdown Message	5-13

This page is intentionally left blank

1. BEFORE YOU START



Warning

- Read this manual to become familiar with all operating procedures and clinical requirements before attempting to operate the system.
- The MARGINPROBE[®] system may be serviced only by Dune Medical Devices qualified personnel.

Conventions Used in this Manual

The following conventions in the form of notes, cautions and warnings are used in this manual:

	Note
Note	

The content of this *Note* offers general information that is important to keep in mind.



Caution

A *Caution* alerts the user to the possibility of a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the user or damage to the equipment.



Warning

A *Warning* alerts the user to the possibility of injury, death, or serious adverse reactions associated with the use or misuse of the system.

This page is intentionally left blank

2. SAFETY AND REGULATORY ISSUES

This chapter describes the safety issues regarding the use and maintenance of the MARGINPROBE[®] system, with special emphasis on electrical safety.

The system is designed for safe and reliable use, when used in accordance with proper operation procedures. The user and all other personnel operating the system should be familiar with the safety information provided in this chapter. Dune Medical assumes no liability whatsoever for any damage or injury as a result of an application of a product which is not in strict accordance with the instructions provided with the product.



Warning

- Read this chapter to be familiar with all of its safety requirements and operating procedures prior to operating the system.
- The MARGINPROBE[®] probe is designed for use only with the MARGINPROBE[®] console.
- High voltage is present inside the console.
- Always be aware of the possible dangers and take proper safeguards as described in this guide.
- For complete contact information please refer to page A of this manual.

Intended Use

The Dune MARGINPROBE[®] System is an adjunctive diagnostic tool for identification of cancerous tissue at the margins (≤ 1 mm) of the main ex-vivo lumpectomy specimen following primary excision and is indicated for intraoperative use, in conjunction with standard methods (such as intraoperative imaging and palpation) in patients undergoing breast lumpectomy surgery for previously diagnosed breast cancer.

Warnings, Precautions, and Contraindications

Contraindications

The Dune MARGINPROBE[®] System should not be used:

- To replace standard tissue histopathology assessment
- On *ex-vivo* lumpectomy specimens that have been exposed to saline, ultrasound gel or local anesthetic solutions.
- On *in-vivo* tissue (i.e. it should not be used within the lumpectomy cavity)
- On tissues other than breast tissue (i.e. it should not be used on Sentinel Lymph Nodes)
- Closer than 1.5 mm to a fine needle localization guidewire

<u>Warnings</u>

- The MARGINPROBE[®] should be used on tissue specimens within 20 minutes of excision.
- The MARGINPROBE[®] should not be used in patients who undergo full cavity excision following removal of the main lumpectomy specimen during the initial lumpectomy procedure.
- The MARGINPROBE[®] has not been studied in patients with:
 - Multicentric disease (histologically diagnosed cancer in two different quadrants of the breast), unless resected in a single specimen
 - Bilateral disease (diagnosed cancer in both breasts)
 - Neoadjuvant systemic therapy
 - Previous radiation in the operated breast
 - Prior surgery at the same site in breast
 - Implants in the operated breast
 - Pregnancy
 - Lactation
 - Cryo-assisted localization

Precautions

- The main *ex-vivo* lumpectomy specimen is defined as the initially excised lumpectomy specimen, without any of the lumpectomy cavity shavings that may have been subsequently taken during the procedure. The device has not been studied for use on tissue shavings excised from the lumpectomy cavity.
- The MARGINPROBE[®] system should be used in addition to standard intraoperative methods of assessing margin status.
- Moving the probe before suction release may potentially damage and affect tissue histopathology
- The MARGINPROBE[®] Probe should only be used with the MARGINPROBE[®] Console.
- The MARGINPROBE[®] Probe is designed for single patient, single-use only and must be properly discarded after use.
- The MARGINPROBE[®] Probe is supplied sterile. If the sterile pack is torn or has been opened, do not use the probe.
- Do not use a MARGINPROBE[®] Probe that has passed its expiration date

Potential Adverse Effects of the Device on Health

Below is a list of the potential adverse effects (e.g., complications) associated with the use of the device.

- Extension of procedure time
- Errors in device reading
- Unnecessary removal of healthy tissue with a potential negative impact on cosmetic results or cosmetic appearance.
- Infection
- Local tissue damage
- Bleeding

For the specific adverse events that occurred in the clinical studies, please see the next section (Clinical Data) below.

Clinical Data

MarginProbe Pivotal Study:

Dune Medical Inc. performed a clinical pivotal study to establish a reasonable assurance of safety and effectiveness of the MarginProbe System. The MarginProbe System is an adjunctive diagnostic tool for identification of cancerous tissue at the margins (≤ 1 mm) of the *ex-vivo* lumpectomy specimen following primary excision and is indicated for intraoperative use, in conjunction with standard methods (such as intraoperative imaging and palpation) in patients undergoing breast lumpectomy surgery for previously diagnosed breast cancer in the US. The pivotal study was performed under IDE # G070182. Data from this clinical study were the basis for the PMA approval decision. A summary of the clinical study is presented below.

A. Study Design

Patients were treated between September 2008 and March 2010.

The MarginProbe System pivotal study was a prospective, multicenter, randomized (1:1), controlled, double-arm study. Breast cancer patients were randomized to either receive standard of care (SOC) lumpectomy or Standard of Care lumpectomy with adjunctive MarginProbe device use (SOC + Device).

Key Aspects of the protocol are as follows:

1. Patient Study Inclusion and Exclusion Criteria

Enrollment in the pivotal study was limited to patients who met the following inclusion criteria:

- Women histologically diagnosed with carcinoma of the breast
- Women with non-palpable malignant lesions, requiring image guided localization.
- Undergoing lumpectomy (partial mastectomy) procedure.
- Age 18 years or more
- Signed informed consent form

Patients were not permitted to enroll in the pivotal study if they met any of the following exclusion criteria:

- Multicentric disease (histologically diagnosed cancer in two different quadrants of the breast)
- Bilateral disease (diagnosed cancer in both breasts)
- Neoadjuvant systemic therapy
- Previous radiation in the operated breast
- Prior surgical procedure in the same breast
- Implants in the operated breast
- Pregnancy
- Lactation

2. Patient Treatment

Patients were first enrolled and taken to the operating room for resection of the main lumpectomy specimen. The main lumpectomy specimen and lumpectomy cavity palpation and related re-excisions were performed before patient randomization. For all main specimens, the center of each of the 6 margins was suture marked. Patient were then randomized to either the SOC or SOC+Device arm intraoperatively, immediately after the main lumpectomy specimen was excised, oriented, center marked, palpated, and additional palpation based re-excision performed.

For patients randomized to the SOC+Device arm the surgeon:

- Applied the MarginProbe device to each of the 6 faces of the excised main lumpectomy specimen—sampling 5 8 points (and up to 12 points for larger specimens). The points sampled were at both evenly spaced and suspicious sites.
- Was required to react to Device feedback. A single positive reading on any margin classified that margin as positive and required the surgeon to remove additional tissue from that margin.
- Documented the reasons why additional margins were not re-exicised despite a positive MarginProbe device reading. For the purposes of CSR primary endpoint

calculations, lumpectomy cavity shavings that were not possible due to physical limitations (proximity to the skin or pectoralis fascia) the margin was considered "addressed".

- Was instructed <u>not</u> to use the MarginProbe device on shavings from the lumpectomy cavity shavings (even if a shaving was taken prior to randomization)
- Was instructed <u>not</u> to use the MarginProbe device within the *in-vivo* lumpectomy cavity.
- Was instructed <u>not</u> the use the MarginProbe device on *ex-vivo* lumpectomy tissue that had been exposed to saline or ultrasound gel. It was however acceptable to use the MarginProbe device on *ex-vivo* lumpectomy tissue exposed to sterile water.
- Was instructed <u>not</u> to use the MarginProbe device in the 1.5 mm region of tissue surrounding a fine needle localization guidewire.

For both SOC and SOC+Device arm patients, lumpectomy specimens were imaged by ultrasound or radiography after randomization and device use. Additional lumpectomy cavity re-excisions were taken as deemed appropriate based on specimen imaging results. Figure 2-1 provides a diagrammatic representation of the study design.

Note that the study design allows for an additional option to perform lumpectomy cavity shavings in the SOC+Device arm (option for shaving at 3 time points) versus the SOC arm (option for shaving at 2 time points).



Figure 2-1: Pivotal Study Design

The MarginProbe device was not used during lumpectomy reoperations.

The study consisted of two phases – a training phase and a randomization phase. Each surgeon had to complete the training phase before being able to randomize patients. Surgeons who had attended 2 or more device procedures (training or randomized) were certified in device use.

3. <u>Pathology Protocol</u>

Pathological assessment was standardized and identical for both study arms. Pathologists were blinded to randomization.

A positive margin was to be defined in this study as a margin microscopically measured and reported in the histopathology report to have cancer within 1 mm or less of the inked surface.

Each investigational site performed the histopathology assessment using a Standard Operating Procedure. Re-cut slides from the first 4 patients at each investigational site (Training, SOC, or SOC+Device) were to be sent to a core-lab and were to be used to review the accuracy and reporting capabilities of the investigational site pathology.

Dimensions (L, W, D) of all excised tissues were recorded. Tissue volume was determined by use of the ellipsoid formula:

 $V = (4/3) * \pi * L * W * D$

4. Duration of Patient Follow-up

Patients were followed until the end of the lumpectomy procedure. Data were collected regarding all ipsilateral breast surgical procedures and their respective permanent histopathology data. Data were to be collected up until the earlier of the following events: conversion to mastectomy, initiation of chemotherapy or two months after the surgery date.

5. <u>Study Endpoints</u>

The prespecified study endpoints are as follows:

Safety evaluation consisted of assessment of all adverse events and serious adverse events, which were summarized using descriptive statistics.

The primary effectiveness endpoint (CSR) is measured as all pathologically positive margins on the main specimen being intraoperatively re-excised or "addressed". A re-

excised or "addressed" margin does not mean that the final true outermost margin is pathologically negative for cancer.

- A positive margin is defined as a margin microscopically measured and reported in the histology report to have cancer within 1 mm or less of the inked margin.
- The main specimen is defined as the lumpectomy specimen removed prior to patient randomization. The main lumpectomy specimen <u>does not</u> include additional shavings even if the cavity shaving was performed prior to patient randomization.
- If a margin has been indicated as positive by the device and documented to not have been re-excised as required by protocol, due to resection already undermining the skin or reaching the pectoralis fascia, this margin will be counted as "detected" and "addressed" for the purpose of CSR endpoint calculation although it was not "re-excised".

An illustration of how CSR is determined is provided in Figure 2-2.

CSR 1º Effectiveness Endpoint

CSR = All positive margins on the main specimen being re-excised/ addressed intraoperatively from positive main specimen cohort (PSS)



Figure 2-2: Illustration of CSR Primary Endpoint

Figure 2-3 below illustrates how the CSR assessment includes both clinically relevant scenario which is the conversion of a specimen which has a pathologically positive for cancer margin to a specimen with negative for cancer margins and the clinically irrelevant scenario in which the additional shaving resulted in the true outermost margin of the specimen remaining pathologically positive for cancer.



Figure 2-3: CSR and Clinical Relevance

While CSR is a focused assessment that is limited to what is within the control of the MarginProbe device, there are limitations to the CSR primary effectiveness endpoint. Some of these limitations are present because the reason and timing for taking additional shavings of the lumpectomy cavity were not documented—that is, whether a shaving was taken because of clinical suspicion, imaging, other assessment, versus a positive MarginProbe device reading and whether the shaving was taken before randomization or after specimen imaging. While the device readings for each margin and the margins shaved were documented, the timing of each shaving and the reason prompting the shaving was not collected.

Table 2-1 summarizes the strengths and limitations of the CSR primary effectiveness endpoint for the pivotal study.

Strengths	Limitations
A focused assessment limited to what is within the control of the MarginProbe device i.e. causing additional cavity shavings.	The study design allows for an additional option to perform cavity shavings in the SOC+Device arm versus the SOC arm. The additional option in the SOC+Device arm may be responsible for an increase in CSR in the SOC+Device arm.
A by specimen assessment which does not give partial credit to intraoperative re- exision of some positive margins on the main specimen but not all positive margins on the main specimen.	The incremental contribution of the MarginProbe device to a higher CSR cannot be determined because the reason for taking a cavity shaving - i.e. SOC (clinical suspicion, or imaging) versus a positive MarginProbe reading - was not documented.
	Questionable clinical relevance. CSR considers whether a shaving was taken or not taken at positive margins on a lumpectomy specimen. CSR does not consider whether the shaving taken converted the initially positive for cancer margin to a negative for cancer final margin.
	CSR does not penalize false positive MarginProbe readings in the positive main specimen cohort. False positive MarginProbe readings in the positive main specimen cohort cause the resection of healthy tissue.
	CSR does not consider false positive MarginProbe readings in the negative main specimen cohort. False positive MarginProbe readings in the negative main specimen cohort cause the resection of healthy tissue.

Table 2-1 - Strengths and limitations of the primary effectiveness endpoint, CSR

Secondary effectiveness endpoints are summarized in Table 2-2 below.

Endpoint	Definition
Incomplete Surgical Re-excision	Proportion of patients with at least 1 positive margin not resected/addressed.
	Differs from primary effectiveness endpoint, CSR, since Yes/No definitions are opposite.
	from the AVS dataset rather than the PSS dataset.
Full Detection	Rate of patients with all positive margins on main specimen detected by device
Re-excision Procedure Rate	Rate of repeated ipsilateral breast surgical procedures (including mastectomies)
Positive Margin Presence	Rate of patients with at least 1 positive margin remaining after lumpectomy
TTV excised in the primary lumpectomy procedure (cm ³)	Average volume of total amount of tissue excised in lumpectomy

 Table 2-2 - Secondary Effectiveness Endpoints

6. <u>Pre-Specified Analysis Plan</u>

For the primary efficacy analysis, a sample size of 116 valid primary effectiveness patients per arm was determined to provide at least 90% power to demonstrate superiority of SOC+Device over SOC.

The analysis populations are defined in Table 2-3.

 Table 2-3 - Analysis Populations

Analysis Population	Definition
All Valid Subjects (AVS)	The AVS subjects included all randomized patients with valid histology data (and valid MarginProbe System data in Device arm)
Positive Specimen Subjects (PSS)	The PSS subject is a subset of the AVS Analysis Set of subjects with at least 1 histologically positive main specimen margin at depth $\leq 1 \text{ mm}$
Negative Specimen Subjects (NSS)	The NSS subject is a subset of the AVS Analysis Set of subjects with no histologically positive main specimen margin at depth ≤ 1 mm.

Safety was assessed using the AVS population. The primary effective endpoint was based on PSS population, and the secondary effectiveness endpoints were based on AVS, PSS or NSS populations as shown in Tables 2-4 and 2-5.

Table 2-4 - The Primary Effectiveness	s Endpoints Population
---------------------------------------	------------------------

Endpoint	Analysis Population	Scoring
CSR	PSS analysis set	Complete Surgical Re-excision (CSR) was scored dichotomously as follows:
		No: At least one positive margin on the main specimen not re-excised/addressed intraoperatively.
		Yes: All positive margins on the main specimen re- excised/addressed intraoperatively

Table 2-5 - The Secondary Effectiveness Populations

Endpoint	Analysis Population	Scoring
Incomplete Surgical Re-excision	AVS analysis set. The groups were compared using 2- sided Fisher's Exact Test.	 Incomplete Surgical Re-excision ("re-excision is used to mean "resection) was scored dichotomously: Yes: If at least 1 positive margin with d ≤ 1 mm on the main specimen was not resected/addressed intraoperatively. No: Otherwise This endpoint differed from the primary effectiveness endpoint, Complete Surgical Resection, since the Yes/No definitions were opposite.
Full Detection	PSS analysis set A 2-sided exact binomial 95% CI for the proportion of "Yes".	Scored dichotomously for SOC+Device arm patients only: Yes: If all positive margins on the main specimen with $d \le 1$ mm were detected by the device (in Device arm) No: Otherwise
Re-excision Procedure Rate	AVS analysis set Compared the groups using a Poisson regression model.	Number of repeated ipsilateral breast surgical procedures (including mastectomies) for each patient. This endpoint was counted as an integer per patient; the count was increased by 1 with each subsequent surgery.

Endpoint	Analysis Population	Scoring
Positive Margin	AVS analysis set	Scored dichotomously.
Presence		
	Compared the	Yes: If there was at least 1 positive margin with $d \le 1$ mm
	groups using a	after the first lumpectomy
	Poisson regression	
	model.	No: Otherwise
TTV excised in	NSS analysis set	Total amount of tissue excised during lumpectomy for
the primary		each patient.
lumpectomy	Compared the	
procedure (cm ³)	groups using a	
	2-sided Wilcoxon	
	Rank-Sum Test.	

The margin-level and patient level (ignoring location) sensitivity and specificity are reported for diagnostic performance of the MarginProbe device. These were not pre-specified in terms of an acceptable minimal sensitivity and specificity. The results here are based on the observed performance in the clinical pivotal study.

B. Subject Accountability

A total of 664 patients who were eligible for study enrollment underwent surgery and were allocated to either the roll-in group or randomization (enrollment allocation). Sixty-eight women were operated on in the roll-in phase and 596 were randomized equally to the Control (SOC arm) and Device treatment (Device +SOC arm) groups. All 664 women completed the study. Subject accountability is displayed below in Table 2-6.

Disposition	Total n (%)
Eligible for Participation	721
Did Not Enter Study	57 (7.9)
Failed eligibility	25 (3.5)
Withdrew consent	6 (0.8)
Other	26 (3.5)
Eligible for Allocation	664 (92.1)
Allocated to Enrollment	664 (100)
Roll-in	68 (10.2)
Randomized to Treatment	596 (89.8)
Device	298 (44.9)
Control	298 (44.9)
Completed Study	664 (100)
Did Not Complete	0 (0)

Table 2-6 - Patient Accountability, Pivotal Study

All 664 women were included in the Safety analysis set. The AVS analysis set includes 596 randomized (298 Device and 298 Control) patients and differs from safety analysis set in 64 roll-in women, as shown in Table 2-7.

		T	Treatment Group		
		Device	Control	Roll-In	Total
Analysis Set	Patients Included	n (%)	n (%)	n (%)	n (%)
Safety Set	All patients for whom	298	298	68	664
-	surgical procedure was	(100.0)	(100.0)	(100.0)	(100.0)
	initiated				
Effectiveness	Sets				
AVS	All Randomized Patients	298	298	NA	596
		(100.0)	(100.0)		(100.0)
PSS	Positive Specimen Patients	163	147	NA	310
	-	(54.7)	(49.3)		(52.0)
NSS	Negative Specimen Patients	135	151	NA	286
		(45.3)	(50.7)		(48.0)

Table 2-7 - Data Sets Analyzed: Number of Patients

All randomized patients completed the study protocol. There was no loss to follow-up in the study. There was no missing data related to the CSR endpoint; 38/1788 (2%) of margins were not measured by the device.

C. Demographics and Baseline Characteristics

Demographic characteristics were similar for the Device and Control groups. Overall, the groups appeared to be comparable, as shown in Table 2-8 and 2-9.

 Table 2-8 - Demographics by Treatment Group

		Treatment Group			
	Roll-In	Device	Control		
Parameter	N=68	N=298	N=298		
Ethnic Origin n (%)					
White ^a	59 (86.8)	250 (83.9)	260 (87.2)		
African-American or Black	5 (7.4)	22 (7.4)	17 (5.7)		
Asian	2 (2.9)	12 (4.0)	10 (3.4)		
Native Hawaiian or Other	0 (0)	3 (1.0)	1 (0.3)		
Pacific Islander					
Other	2 (2.9)	11 (3.7)	10 (3.4)		

^a Includes Hispanics.

		Treatme	ent Group
	Roll-In	Device	Control
Parameter	N=68	N=298	N=298
Age (yrs) Mean (SD)	63.6 (11.1)	60.3 (11.4)	60.2 (11.1)
BMI (mean)	28.0	27.9	28.6
Bra Cup Size n (%)			
AA	0 (0.0)	2 (0.7)	4 (1.3)
А	6 (8.8)	16 (5.4)	16 (5.4)
В	21 (30.9)	101 (33.9)	73 (24.5)
С	24 (35.3)	99 (33.2)	93 (31.2)
D	12 (17.6)	62 (20.8)	92 (30.9)
Е	1 (1.5)	2 (0.7)	5 (1.7)
F	1 (1.5)	1 (0.3)	1 (0.3)
>F	1 (1.5)	1 (0.3)	2 (0.7)
Unknown	2 (2.9)	14 (4.7)	12 (4.0)

 Table 2-9 - Baseline Characteristics by Treatment Group

Table 2-10 presents the number of patients with a diagnosis, requiring that certain categories be combined. For patients with invasive types of carcinoma the mixed invasive category was used, and for patients with more than 1 diagnosis who did not have more than one type of invasive carcinoma, the mixed category was used. The treatment groups appear to be similar with respect to diagnosis.

Patient Diagnosis	Т			
	Device Control		Roll-In Phase	All
	N (%) Patients	N (%) Patients	N (%) Patients	N (%)
Invasive Ductal Carcinoma	24 (8.1)	22 (7.4)	7 (10.3)	53 (8.0)
Invasive Lobular Carcinoma	. 26 (8.7)	13 (4.4)	2 (2.9)	41 (6.2)
Mixed Invasive ^a	8 (2.7)	5 (1.7)	1 (1.5)	14 (2.1)
Ductal Carcinoma in Situ	83 (27.9)	78 (26.2)	19 (27.9)	180 (27.1)
Tubular Carcinoma	1 (0.3)	0 (0.0)	0 (0.0)	1 (0.2)
Mucinous Carcinoma	1 (0.3)	1 (0.3)	0 (0.0)	2 (0.3)
Mixed ^b	155 (52.0)	179 (60.1)	39 (57.4)	373 (56.2)
Total	298 (100.0)	298 (100.0)	68 (100.0)	664 (100.0)

 Table 2-10 - Patient Diagnosis by Treatment Group (Per-diagnosis Analysis)

a Mixed invasive=Invasive Ductal Carcinoma+Invasive Lobular Carcinoma.

b Mixed=more than 1 diagnosis and not only invasive carcinoma.

Tumor stage results are presented in Table 2-11 below. The majority of patients were diagnosed with stage II breast cancer and below.

	()]	[Ι	Ι	I	Π	Γ	V	Unkı	nown	То	tal
Treatment Group	N	%	N	%	N	%	N	%	N	%	N	%	Ν	%
Device	81	27.2	155	52.0	51	17.1	4	1.3	1	0.3	6	2.0	298	100.0
Control	84	28.2	161	54.0	44	14.8	6	2.0	0	0	3	1.0	298	100.0
Roll-In Phase	21	30.9	34	50.0	12	17.6	1	1.5	0	0	0	0	68	100.0
All	186	28.0	350	52.7	107	16.1	11	1.7	1	0.2	9	1.4	664	100.0

Table 2-11 - Tumor Stage

Receptor status is presented in Table 2-12. There were 84 subjects in device and control arms, and 19 in the roll-in subjects, for which HER2 status was not preformed.

Table 2-12 - Receptor Status

Receptor Status	Roll-In N=68	Device N=298	Control N=298
ER+	60/68 (88.2)	251 (84.2)	258(86.6)
PR+	52/68 (76.4)	223 (74.8)	217 (72.8)
HER2+	3/49 (6%)	20/214 (9%)	33/214 (15%)
HER2-	42/49 (85%)	175/214 (82%)	163/214 (76%)

D. Surgical Procedure

The mean duration of anesthesia time (hours: minutes) was 2:03 for the Device group, 1:52 for the Control group and 2:11 for the Roll-in group. This time includes surgical procedures, resections, completion of the protocol procedures, and device use. The mean duration of device use was 5 minutes for the Device group and 6 minutes for the Roll-in group.

Table 2-13 presents the number and percent of patients with a palpable tumor excised during lumpectomy. While all patients had non-palpable lesions at screening (inclusion criteria), the lesion may or may not have been palpable in the *ex-vivo* lumpectomy specimen. There were no apparent differences between treatment groups with respect to palpable tumors during excision.

Was The Tumor Palpable in	Device	Control	Roll-In Phase	All
The Excised Specimen?	N (%) Patients	N (%) Patients	N (%) Patients	N (%) Patients
No	196 (65.8)	188 (63.1)	43 (63.2)	427 (64.3)
Yes	102 (34.2)	110 (36.9)	25 (36.8)	237 (35.7)
Total	298 (100.0)	298 (100.0)	68 (100.0)	664 (100.0)

Table 2-13 - Frequency Distribution of Palpable Tumor during Lumpectomy by Treatment Group

Source: Statistical Table M-38 in Appendix 10.2.2.

Various intraoperative evaluations were used at surgeon discretion in both the SOC and SOC+Device arms and included radiological exam, ultrasound, ultrasonic guidance, touch cytology, gross assessment, and frozen section.

The reason for performing a lumpectomy cavity shaving—that is, whether a shaving was prompted by gross visualization/palpation, positive MarginProbe device readings, imaging, touch prep cytology or frozen section analysis--was not documented.

The methods of excision used during lumpectomy included the following: electrocautery, sharp excision, and scissors.

Table 2-14 describes number of patients undergoing SLNB with dye or radioisotope or both.

 Table 2-14 - Number of Patients undergoing SLNB with Dye or Radioisotope or

 Both

	Roll-In	Device	Control
	N=68	N=298	N=298
SLNB performed	59 (72%)	223 (75%)	225 (75)

E. Pathology

Table 2-15 presents weight and volume of the main specimen. There were no apparent differences between treatment groups with respect to weight and volume of the main specimen. The mean size (diameter) of the main specimen was 4.85 cm for the Device group, 4.89 cm for the Control group, and 4.7 cm for the Roll-in group.

 Table 2-15 - Descriptive Statistics of Specimen Weight and Volume by Treatment

 Group

Specimen	Device		Control		Re	oll-In Phase		All
Parameter	n	Mean (SD)	n	Mean (SD)	n	Mean (SD)	n	Mean (SD)
Weight (g)	294	51.4 (42.2)	290	55.8 (49.8)	67	48.6 (69.4)	651	53.0 (49.0)
Volume (cm ³)	296	59.7 (51.4)	298	61.3 (52.5)	68	54.6 (67.5)	662	59.9 (53.7)
Volume (cm ³)	296	59.7 (51.4)	298	61.3 (52.5)	68	54.6 (67.5)	662	59.9

Source: Statistical Table M-46 in Appendix 10.2.2.

Overall mean tumor size was similar for the groups (MarginProbe=1.7 cm³, Control=1.6 cm³).

The tumor type (as assessed by post-operative histopathology) by treatment group are presented in Table 2-16. The treatment groups appear to be similar with respect to tumor type. The number of positive margins on the main specimen, by treatment group, also appears to be similar.

		Treatment Group					
Tumor Type	Device	Control	Roll-In Phase	All			
	N Specimens (%)	N Specimens (%)	N Specimens (%)	N Specimens (%)			
Invasive ductal	158 (53.0)	179 (60.1)	40 (58.8)	377 (56.8)			
Invasive lobular carcinoma	46 (15.4)	26 (8.7)	9 (13.2)	81 (12.2)			
Ductal carcinoma in-situ	207 (69.5)	229 (76.8)	46 (67.6)	482 (72.6)			
Tubular Carcinoma	5 (1.7)	6 (2.0)	2 (2.9)	13 (2.0)			
Mucinous Carcinoma	10 (3.4)	3 (1.0)	2 (2.9)	15 (2.3)			
Medullary Carcinoma	0 (0.0)	0 (0.0)	0 (0.0)	0 (0.0)			
Papillary Carcinoma	0 (0.0)	2 (0.7)	1 (1.5)	3 (0.5)			
Non malignant (NM)	19 (6.4)	19 (6.4)	5 (7.4)	43 (6.5)			
Other	5 (1.7)	7 (2.3)	0 (0.0)	12 (1.8)			
Total Patients	298 (100.0)	298 (100.0)	68 (100.0)	664 (100.0)			

 Table 2-16 - Frequency Distribution for Tumor Type by Treatment Group

The average weight and volume of resected margins by treatment group during the lumpectomy is presented in Table 2-17. The treatment groups appear to be similar with respect to weight and volume of resected margins.

Table 2-17 - Descriptive Statistics of Resea	ted Margins We	eight and Volume by
Treatment Group		

		Treatment Group						
Specimen		Device Control Roll-In Phase				All		
Parameter	nª	Mean (SD)	nª	Mean (SD)	nª	Mean (SD)	nª	Mean (SD)
Weight (g)	1000	6.6 (6.8)	329	7.5 (6.7)	219	6.0 (5.2)	1548	6.7 (6.6)
Volume (cm ³)	1044	7.9 (10.7)	344	9.1 (10.1)	252	7.4 (8.2)	1640	8.1 (10.2)

^a Difference between weight and volume in number of margins is due to missing data. Source: Statistical Table M-54 in Appendix 10.2.2.

F. Study Results

1. Safety Results

14 adverse events (AEs) were reported, all being categorized as serious adverse events (SAEs) per study protocol definition. One SAE was possibly related to the study device, a wound infection requiring hospitalization and treatment with antibiotics.

Table 2-18 - Frequency of Serious (All) Adverse Events by System Organ Cla	ass,
Preferred Term, and Treatment Group	

		Treatment Group							
		Device N=298		Control N=298		Roll-In Phase N=68		Any N=664	
System	n Organ								
Class/l	Preferred	Ν	N (%)	Ν	N (%)	Ν	N (%)	Ν	N (%)
Term	I	SAEs	Patients	SAEs	Patients	SAEs	Patients	SAEs	Patients
Any	Any	6	6 (2)	5	5 (2)	3	3 (4)	14	14 (2)
	Any	2	2(1)	1	1 (0)	2	2 (3)	5	5 (1)
	Acute tonsillitis	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)
Infections	Breast abscess	0	0 (0)	1	1 (0)	0	0 (0)	1	1 (0)
and	Cellulitis	0	0 (0)	0	0 (0)	1	1 (1)	1	1 (0)
infestations	Postoperative wound								
	infection	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)
	Urinary tract infection	0	0 (0)	0	0 (0)	1	1(1)	1	1 (0)
	Any	2	2 (1)	3	3 (1)	0	0 (0)	5	5 (1)
. .	Fractured sacrum	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)
poisoning and	Post procedural haemorrhage	0	0 (0)	2	2 (1)	0	0 (0)	2	2 (0)
procedural complications	Procedural dizziness	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)
	Procedural pain	0	0 (0)	1	1 (0)	0	0 (0)	1	1 (0)
Neoplasms benign, malignant and	Any								
unspecified		1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)
(incl cysts and polyps)	Uterine leiomyoma	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)

				Treatment Group					
~ ·		Dev		Control		Roll-In Phase		Any	
Systen	n Organ	N=298		N=298		IN=68		N=664	
Class/	Preferred	N	N (%)	N	N (%)	N	N (%)	N	N (%)
Term	Term		Patients	SAEs	Patients	SAEs	Patients	SAEs	Patients
Reproductive	Any	0	0 (0)	1	1 (0)	0	0 (0)	1	1 (0)
system and	Breast								
breast	haematoma								
disorders		0	0 (0)	1	1 (0)	0	0 (0)	1	1 (0)
	Any	1	1 (0)	0	0 (0)	1	1(1)	2	2 (0)
Vascular	Hypertension	0	0 (0)	0	0 (0)	1	1(1)	1	1 (0)
disorders	Hypertensive								
	crisis	1	1 (0)	0	0 (0)	0	0 (0)	1	1 (0)

Adverse events associated with device malfunction or incorrect device readings causing incorrect surgeon action is both a safety and an effectiveness issue. Incorrect surgeon action is therefore further discussed in the Effectiveness Results section below. While an approximately 5 minute prolongation of the operative procedure associated with device use, this prolongation cannot be associated with specific patient adverse events. In addition, while damage to the tissue exposed to the MarginProbe device is a potential problem, an assessment for tissue damage was not considered to be feasible in the pivotal study. From the available data this issue has not been reported.

2. <u>Effectiveness Results</u>

Primary Effectiveness Endpoint: There were a total of 163 patients in the SOC+Device arm and a total of 147 patients in the SOC arm who were in the PSS dataset (i.e. with at least one positive margin by histology on the main specimen). The CSR primary effectiveness endpoint results are provided in Table 2-19.

The device failed to give a reading on 38 (2%) margins out of 1788 margins measured from 298 subjects. This did not impact the primary endpoint.

Primary Endpoint	Dataset	SOC + Device	SOC	Difference (95% CI)	
CSR	PSS	71.8% (117/163)	22.4% (33/147)	49.3% (39.0%,58.7%)	p < 0.0001

Secondary Endpoints	Dataset	SOC + Device	SOC	p-value or CI
Incomplete Surgical Re-excision	AVS	15.4% (46/298)	38.3% (114/298)	p < 0.0001*
Full Detection	PSS	62.6% (102/163)	NA	95% CI: 54.7% - 70%*
Re-excision Procedure Rate	AVS	20.8% (82/298)	25.8% (94/298)	p = 0.3177*
Positive Margin Presence	AVS	30.9% (92/298)	41.6% (124/298)	p = 0.0082*
TTV excised in the primary lumpectomy procedure (cm ³)	NSS	92.7 cm^3	69.9 cm ³	p = 0.0031*

Table 2-20 - Secondary Effectiveness Endpoint Results

* Unadjusted analysis

Of the endpoints listed, the clinically relevant endpoint of re-excision procedure rate showed a 5 percentage point reduction in the SOC+Device arm versus SOC arm.

The reoperation procedure rate is further described in Table 2-21. Note that fewer patients in the SOC+Device arm required a second operation (71 patients in the SOC+Device arm versus 85 patients in the SOC arm). Recall that the MarginProbe device was only used during the initial lumpectomy operation and not during reoperations. More patients in the SOC+Device arm versus the SOC were converted to mastectomy. There are numerous reasons for conversion to mastectomy and therefore this finding cannot be directly attributable to device use.

Table 2-21 - Reoperation Procedure Rate	
Re-excision (including conversion to mastectomy)	

	Lumpectomy	Additional Resections			Total	p-Value
Procedure #	1	2	3	4		
SOC+Device	298	62	7	2	71 (23.8%)	0.3177
SOC	298	77	7	1	85 (28.5%)	
Conversion to mastectomy in device arm = $18/298$ p = 0.46 Conversion to mastectomy in control arm = $13/298$						

_

The following additional analyses, Table 2-22 and Table 2-23, provide information regarding diagnostic performance of the device per margin and per patient (ignoring location).

	Sensitivity(%) (95% CI)‡	Specificity(%) (95% CI) ‡	PPV†(%) (95% CI) ‡	NPV†(%) (95% CI) ‡
SOC+Device	73.8 (68.1,79.4)	45.1 (41.8,48.3)	21.6(20.1,23.1)	89.4(87.2,91.4)
SOC	33.9 (27.5,40.5)	83.4 (81.1,85.7)	29.5(25.1,34.3)	86.0(84.8,87.2)
(SOC+Device)- SOC	39.9(31.4,48.1)	-38.3(-42.4, -34.5)	-7.9(-12.8, -3.4)	3.4 (1.0,5.7)
Device only††	75.2(69.3,80.5)	46.4 (42.6,49.9)	22.3 (20.7,23.8)	90.1 (88.0,92.1)
SOC	33.9 (27.5,40.5)	83.4 (81.1,85.7)	29.5(25.1,34.3)	86.0(84.8,87.2)
Device-SOC	41.3(33.0,49.5)	-37.0(-41.4, -33.0)	-7.2(-12.1,-2.6)	4.1(1.8,6.4)

 Table 2-22 - Diagnostic Performance (per-margin)

[†]PPV and NPV calculated using Bayes theorem on sensitivity and specificity, assuming a common prevalence across the two study arms of 17.0%. [‡]95% Bootstrap percentile intervals.

.

.

†† There were 38 margins with a missing device reading.(6 pathology positive margins and 32 pathology negative margins)

Table 2-23 - Diagnostic I	Performance per pati	ent ignoring location

. -

	Sensitivity(%) 95% CI	Specificity (%) 95% CI	PPV†(%) 95%CI	NPV†(%) 95% CI
SOC+Device	98.8(95.6,99.9)	5.9(2.6,11.3)	53.2(52.1,54.4)	81.9(49.0,95.4)
SOC	68.7(60.1,76.1)	53.6(45.4,61.8)	61.6(56.7,66.3)	61.3(54.4,67.7)
(SOC+Device)- SOC	30.1(22.6,38.2)	-47.7(-56.6, -38.3)	-8.4(-13.6, -3.5)‡	20.6(-9.2,42.0)‡
Device only	96.3(92.2,98.6)	8.9(4.7,15.0)	53.4(51.9,54.9)	68.9(46.2,85.2)
SOC	68.7(60.1,76.1)	53.6(45.4,61.8)	61.6(56.7,66.3)	61.3(54.4,67.7)
Device-SOC	27.6%(19.6,36.0)	-44.7% (-54.0, -34.9)	-8.2 (-13.5,-3.1)‡	7.6(-16.6,27.9)‡

[†]PPV and NPV calculated using Bayes theorem assuming a common prevalence across the two study arms of 52%.

\$95% Bootstrap percentile intervals.

The Figures 2-3 and 2-4 provide a more comprehensive assessment of what occurred in each arm of pivotal study.

As shown in Figure 2-4, 298 SOC patients were enrolled. An average of 72 cm3 of tissue was excised during the initial lumpectomy. There were 147 patients with cancer positive main specimens and 151 cancer negative main specimens. Of the 147 cancer positive main specimens, 25 or 17% were converted to cancer negative final margins with cavity shavings.



In the SOC arm, shavings were not taken in 46+81 or 127/298 subjects.

Figure 2-4: Pivotal Study Patient Flow Chart - SOC Arm

As demonstrated in Figure 2-5 298 patients were enrolled in the SOC+Device arm. An average of 88 cm3 of tissue was excised during the initial lumpectomy. There were 163 patients with cancer positive main specimens and 135 cancer negative main specimens. Of the 163 cancer positive main specimens, 79 or 49% were converted to cancer negative final margins with cavity shavings.

In the SOC+Device arm, shavings were not taken in 2+8 or 10/298 subjects.



Figure 2-5: Pivotal Study Patient Flow Chart - SOC+Device Arm

Summary of Supplemental Clinical Information

A. Pivotal Study Additional Analyses

While not powered to detect differences across subpopulations, there was a trend for outside of US patient populations to experience greater clinically relevant benefit than for the US population of patients enrolled as shown in Table 2-24.

		US Patients n = 566		Israel Patients n = 98			
Endpoint		SOC + Device	SOC	SOC + Device	SOC		
1°	CSR	69.7%	22.4%	85.7%	22.7%		
2°	Incomplete Surgical Re-excision	17.3%	38.8%	6.1%	35.4%		
2°	Full Detection*	59.9%	N/A	81%	N/A		
2°	Re-excision Procedure Rate	34.5%	48%	4.8%	22.7%		
2°	Positive Margin Presence	53.5%	82.4%	38.1%	86.4%		
2°	Total Tissue Volume Excised (cm ³)	92.4	82.6	97.6	95.9		
Diagnostic Device Performance		SOC + Device	SOC	SOC + Device	SOC		
Sensitivity (%) 95% CI†		73.4 (66.8,79.6)		87.8 (76.8,98.8)			
Specificity (%) 95% CI†		44.7% (40.8,48.8)		53.9% (46.0,62.0)			

Table 2-24 -	Pivotal	Study	Results	across	Subno	nulations
1 abit 2-2-	IIVUtai	Study	INCSUILS	aci 055	Suppo	pulations

*Full detection is for Device (not SOC+Device arm) †95% Bootstrap percentile intervals.

B. Product Development Clinical Studies

Product development clinical studies were conducted at various stages of the product development process, as summarized in Table 2-25. None of these studies were pre-approved by FDA.

Study	Study Name	#	Product	Primary Objective	Principal Results
Number	Study Mame	[#] Subjects	Description	Trimary Objective	T meipai Results
III	"Point-by-point" study in pathology - phase II 3/2006 – 6/2007	N=76	MarginProbe System Probe & MarginProbe System Type 1.0 system console	Obtain database set and assess performance – phase II	Device use has no permanent effect on tissue (macroscopic or microscopic) Device performance per-point on bread-loafed lumpectomy specimens: sensitivity 100% and specificity 87% on homogeneous samples, sensitivity 70% and specificity 70% on full dataset
V	Intraoperative blinded study - phase II 6/2006 – 5/2008	N=175	MarginProbe System Probe & MarginProbe System Type 1.0 system console	Assess intraoperative performance on the resection surface of lumpectomy specimens and evaluate adjunctive device contribution to SOC	Even with a limited point sampling by the device, per- patient detection rate is superior with Device+SOC (73%) as compared to SOC alone (46%)
MAST	Pilot Study 11/2006 – 11/2007	N=300	MarginProbe System Probe & MarginProbe System Type 1.0 system console	Assessment of device detection performance and clinical utility in a randomized, controlled (patient is blinded), intended use fashion. Assess cosmetic outcome associated with device use compared to SOC.	 Device is safe for intraoperative use Re-excision rate is reduced by 56% (p=0.0027) Positive margin identification guiding intraoperative resection is superior in Device+SOC arm (60%) compared to SOC (41%) Cosmesis is not affected by device use Excised tissue volume is not affected by device Performance is the same for both palpable and non-palpable lesions

Table 2-25 - Summary of Developmental Clinical Studies

The product development study results were used to develop the MarginProbe System algorithm in the manner described in Figure 2-6.



Figure 2-6: Algorithm Development Process

1. Study III

Study III was conducted to create the classification database of actual tissue measurements using the MarginProbe paired with their histology at point level. For each point measured with the device the pathology was taken at that same point. Device measurements were performed at the interior of the lumpectomy specimen (following its sectioning at the pathology lab).

The specimens used for this study were taken from women with palpable tumors who had undergone lumpectomy or mastectomy. The study was performed in Israel at 4 study sites. The patient demographics and cancer specifics of the specimens used to create the classification dataset are summarized in Table 2-26. Table 2-27 illustrates the classification data set that was derived in Study III.
Sites	4 (Israel)	
Ν	77 patients and 81 specimens (4 patients bilateral disease)		
Mean Age (range)	62.64 years (3	6 - 85)	
Mean Tumor Size (range)	1.65 cm (0.1	- 3.5)	
Fine Needle Localization	33 specimens		
Sentinel Node Biopsy (Both Blue Dye & Radioisotope)	43 specime	ens	
Cancer Pathology	Infiltrating Ductal (IDC)	46	
	DCIS	8	
	Mixed	8	
	Infiltrating Lobular (ILC)	6	
	Other	3	
	Not stated	4	
Grade	I	3	
	II	34	
	III	20	
HER2 positive	18		
Estrogen Positive	60		
Progesterone Positive	46		

Table 2-26: Study III - Patient Demographics and Cancer Specifics

Number of tissue measurement data points	869
- Excluded data points	116
Valid data points	753
- Normal	588 (78%)
- Malignant	165 (22%)

 Table 2-27: Study III - Classification Data Set

The ROC curves of the device performance in Study III are shown in Figure 2-7. This figure includes three datasets: (1) tissues containing at least 75% of a single tissue type; (2) all tissues containing at least 50% of a single tissue type; and (3) the full dataset collected in the experiment, containing cancers of all sizes (down to 0.15-mm-diameter features).



Figure 2-7: Study III - ROC curves of 3 different datasets

When the composition of the tissue being measured by the probe (i.e. directly underneath the 7 mm footprint of the probe) was more homogeneous, there was greater sensitivity and specificity in MarginProbe readings as shown in Table 2-28.

Percentage single tissue type within probe's 7 mm diameter footprint	Specimen description	Device Performance
> 75% singe tissue type	22 cancerous, from 15 patients 425 nonmalignant	Sensitivity 1.00 (95% CI: 0.85–1)
		Specificity 0.87 (95% CI: 0.83–0.90)
\geq 50% single tissue type	29 cancerous, from 18 patients, and 567 nonmalignant	Sensitivity 1.00 (95% CI: 0.88–1) Specificity 0.72 (95% CI: 0.68–0.76).
Full dataset containing cancers of all sizes (down to 0.15-mm-diameter features)	165 cancerous sites from 50 patients, and 588 nonmalignant sites	Sensitivity 0.70 (95% CI: 0.63–0.77), Specificity 0.70 (95% CI: 0.67–0.74)

Table 2-28 - Study III - Sensitivity and Specificity in MarginProbe Readings

The performance for different histopathology types are also summarized in Table 2-29. [The two most common groups, invasive ductal carcinoma (IDC) and ductal carcinoma in situ (DCIS), have sensitivities of 0.68 (95% CI: 57–77) and 0.63 (95% CI:45–79), respectively]

Cancer histopathology	Number of samples	Detected	Detection rate (95% CI)
Infiltrating Ductal Carcinoma (IDC)	87	59	0.68 95% CI:57– 77
Ductal Carcinoma in-situ (DCIS)	35	22	0.63 95% CI:45–79
Infiltrating Lobular Carcinoma (ILC)	7	5	0.71
IDC+ DCIS	25	21	0.84
ILC+ DCIS	3	3	1.00
Other	8	6	0.75
Full dataset	165	116	0.70

 Table 2-29: Study III - Device Sensitivity for Different Histopathology Subgroups

2. <u>Study V</u>

Study V was a blinded study with MarginProbe System Type 1.0 device to assess performance of the device on the cut surface tissue of lumpectomy specimens, as compared to histology.

Surgeons were blinded to the device outputs and could not act on device outputs. The device measurements (maximum of 20) were taken intraoperatively on the surface of fresh intact lumpectomy specimens. The orientation of each measurement site was noted. For each marked site, the corresponding 7 mm wide tissue specimen was processed *en-face* and microscopically evaluated as positive or negative for malignancy.



Figure 2-8: Study V - Sampling Process

A total of 175 subjects were enrolled in 3 sites during this study. Surgeons at 2 institutions included in this study (site 1: US site, n=101 patients; site 2: OUS site, n=9 patients) excised additional margins only where deemed necessary ("selective" re-excision). Practice at the third institution (US site, n=65 patients, 66 specimens) was to routinely re-excise all margins from the cavity ("total" re-excision).

While results from Study V served to further inform the MarginProbe product development, Study V also serves to provide a comparison of differences in standard of care selective versus empiric total cavity shaving. Patients who receive empiric, routine re-excision of all margins have greater conversion of initial positive lumpectomy margins to final negative margins. The observed effect is illustrated below in Figures 2-9 and 2-10 comparing the final pathologies from patients treated at study sites 1 and 2 (selective re-excision) versus study site 3 (total re-excision).

There is also literature (see references list below) suggesting that the standard, empiric practice of complete/partial lumpectomy cavity shavings in the same operative setting as the initial lumpectomy can reduce the incidence of incomplete cancer resection and produces greater volumes of tissue resection.



Figure 2-9: Study V - Final Pathologies from Patients Treated at Study Sites 1 and 2 (Selective Reexcision)



Figure 2-10: Study V - Final Pathologies from Patients Treated at Study Site 3 (Total Re-excision)

3. MAST Study

This MAST pilot study was performed in Israel. It was a prospective, randomized, controlled study designed to compared SOC lumpectomy with to SOC+Device lumpectomy. Three hundred subjects at 11 sites were enrolled (n=149 device arm; n=151 control arm).

The MAST study design was similar to the Pivotal study however there were some differences. The MAST study involved a different MarginProbe device algorithm, different device use instructions (i.e. surgeons used the device at their discretion with respect to extent of device use and tissue targeted and were not required to act on positive MarginProbe device readings), an assessment of post-lumpectomy breast symmetry using a 4 point scale, and intra-operative pathology as part of SOC--being used in approximately 20% of the cases.

The difference in protocols across studies may be reflected in the results of the SOC arm in the MAST Study compared to the pivotal IDE investigation. The results are provided in Figures 2-11 and 2-12 below.



Figure 2-11: MAST Study - Final Pathologies - SOC Arm



Figure 2-12: MAST Study - Final Pathologies - SOC+Device Arm

Electrical and Mechanical Safety Precautions

- There are no user-serviceable parts inside the system.
- Make sure that all personnel are familiar with the system's controls and know how to shut down the system instantly in case of an emergency.
- Keep all covers and panels of the system closed. Removing the covers creates a safety hazard.
- Any handling of the system other than intraoperative use should be performed when the system is shut down and disconnected from its electrical power source.
- Move the system slowly and carefully. The system weighs approximately 82 kg (180 lbs.) and may cause injury if proper care is not used.
- The system is grounded through the grounding conductor in the power cable. This protective grounding is essential for safe operation.
- Portable and mobile communications equipment and OR electrical equipment can affect medical electrical devices. Interference may occur in the vicinity of the equipment.
- Do not use any accessories or cables other than those specified and provided. Use of such accessories or cables may result in damage to the system. Connecting any third-party equipment to the system is strictly forbidden without written approval from Dune Medical.

Compliance with International Regulatory Standards

The MARGINPROBE[®] system complies with the requirements of the following:

- IEC 60601-1:1990 + Am. A1+A2: Medical Electrical Equipment, Part 1: General Requirements for Safety. Collateral Standard: Safety Requirements for Medical Electrical Systems
- IEC 60601-1-1:2001: Medical Electrical Equipment Part 1-1: General Requirements for Safety Collateral Standard: Safety Requirements for Medical Electrical Systems
- IEC 60601-1-2:2001: Medical Electrical Equipment, Part 1-2: General Requirements for Safety. Collateral Standard: Electromagnetic Compatibility, Requirements And Tests
- IEC 60601-1-4:1996 + Am. A1: Medical Electrical Equipment -- Part 1-4: General Requirements for Safety. Collateral Standard: Programmable Electrical Medical Systems
- IEC 60601-1-6:2004: Medical Electrical Equipment Part 1-6: General Requirements for Basic Safety and Essential Performance Collateral Standard: Usability
- ISO 10993-1:2003 Biological Evaluation of Medical Devices Part 1: Evaluation and Testing
- ISO 10993-5:1999 Biological Evaluation of Medical Devices Part 5: Test for *in vitro* Cytotoxicity
- ISO 10993-7:2008 Biological Evaluation of Medical Devices Part 7: ETO Sterilization Residuals
- ISO 10993-10:2002 Biological Evaluation of Medical Devices Part 10: Tests for Irritation and Delayed-Type Hypersensitivity
- ISO 10993-11:2006 Biological Evaluation of Medical Devices Part 11: Tests for Systemic Toxicity
- ISO 11135-1:2007 Sterilization of Health Care Products. Ethylene Oxide Requirements for Development, Validation and Routine Control of a Sterilization Process for Medical Devices

Device Regulatory Labels

The following labels are affixed to the system and accessories:

Probe Identification Label

The probe identification label, located on the sealed pack, includes the following information:

- Name and part number of the probe
- Name and address of the manufacturer
- Serial number, lot number, date of manufacture & date of expiration
- Cautionary symbol: Single-use only
- Cautionary symbol: Consult operating instructions
- Cautionary symbol: Do not use if sealed pack is compromised
- Indicator that the probe has been sterilized by ethylene oxide

Console Identification Label

The console identification label, located on the console's rear panel, includes the following information:

- Name and address of the manufacturer
- Name and model number and part number of the console
- Date of manufacture and Serial number of the console
- Electrical rating and fuse type definition
- Cautionary symbol: Follow operating instructions

Electrical Hazard Label

This label warns against dangerous electrical voltages within the console.

EtO Exposure Indicator

This indicator, found on the blister packaging of the probe, is a chemically-sensitive indicator that turns green when exposed to ethylene oxide and indicates that the contents of the pack are sterile. If the indicator on the probe's sealed pack is any other color than green, or if the indicator is missing, do not use the probe!

This page is intentionally left blank

3. CONSOLE PLACEMENT

Equipment List

The MARGINPROBE[®] console shipment package contains the following:

Equipment Item	Quantity
	1
USER Manual – English (this document)	1

Facility Requirements

Before unpacking the system, ensure that the site meets the requirements described in the following sections.

Electrical Requirements

The system will require a separate line supply for the following (appropriate to local requirements):

• 230VAC/50Hz or 115VAC/60Hz, single phase

The electrical requirements are printed on the console identification label (located on the console's rear panel).



Warning

To ensure safe operation, the system must be connected to a properly grounded electrical wall socket.

Space Requirements

Space should be allocated with adequate ventilation and free airflow. The working area for the system should be prepared according to the system dimensions presented in Figure 3-1.



Figure 3-1: System Dimensions

 Caution

 Do not cover the console when switched on.

Operating Environment	General Characteristics
Temperature	15°C to 30°C / 59°F to 86°F
Humidity	30% to 80% at temperature < 29°C / 84°F
Altitude	0 to 2000m (0 to 6560 feet)
Vibration	0.5G maximum, 5Hz to 500HZ

Storage Requirements – MARGINPROBE[®] Console

Storage Environment	General Characteristics
Temperature	10°C to 40°C / 50°F to 104°F
Humidity	20% to 85% RH at temperature < 40°C / 104°F
Altitude	0 to 4572m (0 to 15000 feet)
Vibration	0.5G maximum, 5Hz to 500Hz

Storage Requirements – MARGINPROBE[®] Probe

- Use before the expiry date indicated on the label.
- Store at room temperature below 40°C / 104°F, in a dry place, protected from light.

Console Placement

Console Placement is carried out after uncrating the system from its crate to verify the system is ready for use in a surgical environment and it is performed by doing the following:

- *1.* Unpack the console from its shipping crate.
- 2. Check the console for external damage.
- 3. Verify that the power cable is not damaged and matches the electrical wall socket, connect it and ensure that it is firmly connected to the wall socket.



- 4. Plug the console's power cable into an appropriate electrical outlet.
- 5. Turn on the main power switch (Refer to Figure 3-2):



Figure 3-2: Electrical Connection

- 6. Turn on the console and wait for the "connect probe" message to be displayed on the screen. Instructions can be found in chapter 5, Operating Instruction, Preparing the System for Operation.
- 7. The console is now ready for use.
- 8. Turn off the console as described in chapter 5 System Shut-Down.



4. System Description

Introduction

The MARGINPROBE[®] system consists of a console and a sterile, disposable probe. The MARGINPROBE[®] console consists of internal electrical components, a pneumatic system and a graphic user interface module. The user interface module incorporates a color display, audio components and operation buttons.



Figure 4-1: MARGINPROBE[®] System

Console

The console (Figure 4-2) incorporates modules for signal generation and collection, electric and pneumatic control, data processing and display. The console includes a connector for attaching the probe cable.



Figure 4-2: System Description: Console

Control Panel

The control panel on the front of the console incorporates all system controls as noted in Figure 4-3. These controls will be described in detail in the Operating Instructions chapter (section 5, p 5-1).

Several of the functions performed from this panel may also be performed from the probe; these will be explained in the Operating Instructions chapter.



Figure 4-3: System Description: Control Panel

Summary of control panel buttons

A full description of control panel buttons can be found in the chapter (page 5-1)

User Interface

The user interface (Figure 4-4) consists of an LCD screen and displays the system's status at any given point and the results of the measurements.

The user interface will be described in detail in the Operating Instructions chapter (section 5).



Figure 4-4: Graphic User Interface

Probe Components

The probe is a hand-held, sterile, disposable unit (Figure 4-5) with a 3-year shelf life. It includes the sensor/FFS module, attachment mechanism, calibration module, control button, LED indicators and biological air filter. It is connected to the console with a built-in connector that combines electrical and vacuum connections. The energy is electromagnetic at the RF range. It is confined to the vicinity of the probe tip. The energy level per measurement is less than 0.2 mJ with a power lower than 0.3 mW. The max field voltage is 1V p-p.



Figure 4-5: System Description: Probe

Coupling and Measurement

The tip of the probe incorporates the tissue-sensor attachment mechanism and the sensor that measures the tissue.



Figure 4-6: Probe Tip

Probe Control Button

The system can be operated during a procedure by pressing the control button (Figure 4-7):

- **Double-Click:** for opening and closing measurement frames, refer to Operating Instructions (section 5, p 5-6).
- **Single-Click:** for designating the measurement frames, refer to Operating Instructions (section 5, p 5-6).
- Long Click: for editing and confirming groups of measurement points, refer to Operating Instructions (section 5, p 5-7).



Figure 4-7: Control Button and Probe LEDs

Probe LEDs

The probe features three LEDs (Figure 4-5 and Figure 4-7), each one signifying a different result of a point measurement; during calibration and in standby mode, all the LEDs will be turned on. Refer to the Operating Instructions chapter for complete details (section 5).

Probe Connector

The probe connector (Figure 4-8 and Figure 5-3) couples the probe to the MARGINPROBE[®] console.

The body of the connector and the connection port are notched to each other with a dimple on the inner rim of the connector and a depression for the dimple in the rim of the port, so that the connector may only be inserted in the correct orientation.

A half-turn to the right locks the connector in place.



Figure 4-8: Probe Connector

This page is intentionally left blank

5. OPERATING INSTRUCTIONS

Introduction

The probe was designed to measure breast tissue. Before performing a measurement on the specimen, make sure that the specimen is wiped free of excess fluid. Do not apply saline or ultrasound gel to the tissue before measurements are taken. If necessary, clean the surface of the tissue with sterile water.

An individual measurement is automatically performed by applying the probe tip perpendicular to the tissue and ensuring stable contact for the suction holes in the perimeter. Once in contact with tissue, the tip is automatically attached by slight suction and a measurement is taken.



Warning

- Do not apply saline or ultrasound gel to the tissue before performing a measurement.
- Take measurements within 20 minutes after specimen excision.

Preparing the System for Operation

Caution

- Check the integrity of the screen at the start of every day's scheduled procedures. Do not use the system if the screen is damaged.
- To prevent the system from tipping, ensure that the wheel brakes are **not** locked when the system is in use. Wheel brakes should only be engaged for storage of the console.

The MARGINPROBE[®] system should be turned on and initialized prior to connection of the probe.



The power LED will be orange when the console is plugged in and turned off. Press the start/shutdown switch on the control panel; the power LED will turn green and the system will initialize its software while exhibiting a splash screen with the Dune Medical logo and an initialization progress bar. Typical initialization time is around 3 minutes.

When initialization is complete – and a probe is not connected to the system – a **Connect Probe** message will pop up on the screen (Figure 5-1):



Figure 5-1: Connect Probe Popup Window

The first screen that appears after initialization reflects the way the console was shut down in the previous session. In case of a normal shutdown, a clean screen (without measurements) will appear. If the system was not shut down properly, the screen will display all of the measurements from the previous session. You can always toggle between a new screen and that of the previous session by a long press on the **[T]** button on the control panel.

If no action was taken for ten minutes, the system will switch to **Standby** mode in which a screen saver will appear stating **System in Standby Mode** (Figure 5-2).



Note

- Any popup displayed on the screen can be removed for ten seconds by pressing any of the buttons on the control panel or by a single click on the probe button.
- Resuming operation from **Standby** mode is done by pressing any of the buttons on the control panel or by a single click on the probe button.



Figure 5-2: Standby Mode Message

Connect the Probe outside the Sterile Field

The length of the probe cable is 2.7 meters (8 feet 10 inches) to enable proper maintenance of the sterile field around the patient.

- 1. Circulating Nurse: open the probe's box and remove the plastic blister from the box.
- 2. Circulating Nurse: open the plastic blister outside the sterile field.
- 3. Scrub Nurse: remove the sterile probe from the blister, and hand the connector out of the sterile field.
- **4.** Circulating Nurse: insert the probe connector into the system's connection port. Turn the connector one half turn to the right (clockwise), until you feel it reaches its position (Figure 5-3).

Note	Note
	The connector and the port are both notched to prevent incorrect insertion.



Caution

- Although the console's connector is covered, visually check it to ensure that there are no foreign objects in the connector every time before connecting a new probe.
- Do not use a new probe if the package is damaged. If the package was dropped, visually inspect the probe and do not use if damaged.



Figure 5-3: Probe Connection

Operating the System

The system senses that a probe has been connected and automatically proceeds to the calibration process, during which a **Calibrating** message and progress bar appear at the bottom of the screen (Figure 5-4).

The calibration process takes approximately 10 seconds, during which all the LEDs are illuminated. In the event of a calibration failure a popup message will appear on the screen asking to press the probe's button while holding the probe in the air. A second calibration failure will require replacement of the probe.



Figure 5-4: Calibrating Message

If the system is restoring the operation of a used probe, the same calibrating message will appear on the screen.

When this occurs, the calibrating message will appear and all the LEDs will turn on.

Resuming operation after **Standby** mode will also trigger calibration.

Probe Identification

After the calibration process, the probe type is determined by its ID microchip. The system will be ready to measure the specimen immediately after calibration (Figure 5-5):



Figure 5-5: New Probe Screen

You should connect a used probe only when restoring an ongoing procedure (e.g. if the system was turned off and had to be restarted in the middle of an operation ,such as in the event of a power failure, etc.)



Note

- When connecting a probe to the console, the system identifies the probe, and if the probe was used for a prior procedure an error message is displayed on the screen declaring the probe has exceeded its usage limit.
- Please refer to Table 7-1: "Error message troubleshooting guide" for a complete description of the messages displayed by the console.

T

The first displayed screen depends on how the system was shut down in the previous session. If the system was not shut down properly, the first screen will show the results of the previous session. In all other cases the first screen will be blank. To reach the desired screen to begin measuring press the **T** button for 3 seconds. This will toggle between a clear screen and the screen showing the results of the previous session.

/ Note

Note

- Ensure no previous records are displayed on the screen at the beginning of new procedure.
- By this point during surgery, the surgeon has already removed and blotted the surgical specimen, marked it for orientation, and established the borders & faces of the specimen: Lateral, Superior, Medial, Inferior, Anterior, and Posterior (Deep)

Grouping Measurements According to Margins

Point measurements per margin are grouped together within brackets which facilitates ease of review/analysis. The brackets may be opened and closed using the control button on the probe or the buttons on the control panel:

Double-clicking on the probe control button or pressing the [] button on the control

panel opens a bracket.

- Double-clicking again on the probe button after approving the group (or pressing the [] button on the control panel) closes the bracket.
- A pop up window will appear on the screen with the margin name in red (Figure 5-6).



Figure 5-6: choosing the Margin to be measured.

Single-click the control button to change the margin name and continue to single-click until desired margin label is reached. A bracket with a **red** letter above it, indicating the margin name will also appear on the screen. The letters correspond to the margins as follows:

- L = Lateral
 M = Medial
 I = Inferior
- A = Anterior D = Posterior (Deep)

The margin names may also be changed by pressing the \blacktriangle or \checkmark buttons on the control panel and scrolling to the desired margin label.

After choosing a margin name, press and hold the probe control button to confirm the setting. The letter above the bracket will turn yellow (Figure 5-7) and measurements may be performed. Margin confirmation may also be done from the console control panel by pressing the **T** button.





Note

Note/

- Confirmation of margin label will take place automatically when measuring starts, even if the control button was not pressed.
- To cancel measurement grouping in labeled brackets (before obtaining any measurements), first press and hold the button to approve the name, and then double-click to cancel the brackets.

Grouping Measurements by Numbers: groups may also be named by numbers instead of letters (e.g. for measuring points on the specimen outside of the margin grouping framework). To open a numbered group, scroll to the numbers by pressing the probe control button or the \blacktriangle or \checkmark buttons on the control panel.

Every time a group of measurements is closed, the next numbered group created will be labeled by the next number.

Changing Frame Label: can only be done before closing the brackets. Changing the label is done by pressing and holding the probe control button (to enable editing), and scrolling by clicking to the desired label (or by using the \blacktriangle or \triangledown buttons on the control panel). This can be done even if measurements were taken. After the brackets have been closed, changing the label is no longer possible.

Measurements can also be taken without grouping them by brackets.

	Note
ote	In order to standardize device output and establish a routine protocol, it is recommended to take measurements according to labeled specimen margins.

Performing Measurements

Each individual measurement produces a binary positive/negative display on the screen, as well as an audio indication.

The volume of the audio indication may be adjusted by pressing the up/down volume button on the control panel. When pressing the up/down volume button a beep will be heard. The volume of the beep corresponds to the volume of the audio indication.

When the system is ready for measurement, a partial vacuum is activated to aid in the initial coupling of the probe to the tissue surface. Place the tip of the probe in contact with the tissue point to be measured, and ensure that the entire tip is in contact with the tissue (Figure 5-8). A measurement is automatically initiated when tissue-to-sensor coupling is achieved.

It is recommended to perform 5 to 8 measurements per margin (up to 12 measurements on a large specimen). In each measurement an area of 7 mm in diameter is sampled to the depth of at least 1 mm.



The specimen in Figure 5-8 is an artificial model for illustration purposes.



Figure 5-8: Probe Tip Application to the Specimen

At the end of the measurement the vacuum is released and the probe can easily be removed from the tissue. The table below summarizes possible system outputs:

Measured Indication	LED Color	System Tone	Screen Reaction
Positive (malignant tissue detected)	Red	Single Beep	Red bar above line-RED
Negative (no malignant tissue detected)	Blue	Double Beep	Blue bar below 🔲 line-BLUE
Failed Measurement	Yellow	Error Tone	Empty Bar

Table :	5-1: F	Possible	System	Measurement	Outputs	(Figure	5-9):
1 1000		0000000	<i>System</i>		o mp ms	1 18 11 1	<i>v > j</i> .

Lift the probe after each system tone sounds. If the probe is left on the tissue after the tone, an additional measurement will be taken automatically.



Caution

- Lift the probe only after the tone sounds in order to prevent tearing of the tissue.
- Attach the probe tip only to tissue and avoid direct contact with other materials or surgical tools.

Attempt to evenly sample each margin surface area. In addition, suspect areas may be sampled as deemed necessary.

In case of failed measurements indicated by a blank bar, repeat the measurement.

As mentioned above, the probe is designed for single use, can only be connected to one console and can obtain a limited number of measurements. There is a time limitation for using the probe once it has been connected to the console. When 30 or fewer measurements remain, a counter will appear on the screen indicating the number of measurements left.

Note

Note

If eight consecutive failed measurements are obtained, a message instructing the user to replace the probe will appear on the screen.

Caution

If the probe tip is not lifted from the tissue after a measurement, the system will continue to measure the same point repeatedly.

- Check the probe tip after measuring each margin;
 - $\Rightarrow\,$ If tissue remnants are found, measure the margin again after wiping the probe tip.
- If measurements are taken by the probe without being coupled to tissue or if the sensor plate is found to be jammed a message instructing the user to replace the probe will appear on the screen.
- The probe is for single use only and must be disposed after each case.

Interpreting Measurement Results

The device provides audible and visual indications regarding the assessment of tissue at each point measured.

Visual indication is by a colored bar on the screen that indicates a positive (red) or negative (blue) result. Point and margin data accumulate on the screen.

Using the probe control button, groups of points can be marked and oriented. A margin with one or more positive points is indicated as a "device-positive margin" and should be shaved from the cavity. Device output is adjunctive to other information available intra-operatively to guide the procedure.

The visual results are shown on the screen (Figure 5-9).

- The red bars indicate a positive margin while the blue bars indicate a negative margin.
- The yellow brackets define a group and the letter or number above the bracket represents the group's label.
- The counter on the bottom-left side of the screen indicates the number of measurements that were taken within the current group (bracketed number) and the total number of measurements taken from beginning of the session.



Figure 5-9: Measurement Results

Screen Scrolling

Data points accumulate on the screen from left to right and top to bottom.

When the screen is full, the rows will scroll upwards and data will continue to accumulate in the same manner.

Up to four rows will be displayed on the screen at any given time. Once the screen is full with measurements, the scrolling buttons will appear in the bottom-right corner of the screen. To view hidden rows, scroll up or down by pressing the \blacktriangle or \checkmark buttons on the control panel (Figure 5-10).



Figure 5-10: Screen Scrolling

Probe Usage Limitations

The MARGINPROBE[®] probe is a highly sensitive instrument, designed to detect extremely small differences in tissue characteristics. In order to provide results as intended, it is critical that the sensor be used in the manner validated during clinical development. The performance of the device depends on carefully controlled manufacturing, sterilization, and calibration procedures. The probe is designed and manufactured for single use. The performance of the device cannot be guaranteed if not used in accordance with the instructions provided.

Clearing the Screen and Removing Popup Messages



The screen can be cleared at any point. Press the **T** button to toggle between a clear screen and a screen showing the results up to the last measurement.

Any pop-up displayed on the screen can be removed for ten seconds by pressing the \blacktriangle or \checkmark buttons on the control panel or by single-clicking the probe control button.

Standby Mode

After ten minutes of inactivity, the system will go into **Standby mode** and display a screen saver message: "**System in Standby mode**. **Press any button to resume operation**". Pressing any button on the control panel or the probe will clear the Standby screen message and set the system back to operational (Ready) mode. During standby mode, the pneumatic vacuum module will shut down.

A calibration process will automatically take place when resuming operation after **Standby mode**.

System Shut-Down

To shut down the MARGINPROBE[®] system:



Press the start/shutdown switch on the control panel for three seconds; the system will start the shutdown process and a pop-up message will appear stating: "**System preparing to shutdown. Press Power button to cancel**" (Figure 5-11). After several seconds the system will power down and the power LED will turn orange.

In order to cancel the shutdown process, press the start/shutdown switch again. Pressing the start/shutdown switch for less than three seconds will not initiate the shutdown process.

Turn off the main power switch on the lower-rear panel after shutdown process is complete.



Figure 5-11: Shutdown Message

This page is intentionally left blank
6. MAINTENANCE

Introduction

The MARGINPROBE[®] system is designed to operate reliably without any need for operator maintenance. However, the outer surfaces of the system should be kept clean and free of dust.

Warning

System cleaning should be performed only when the system is shut down and disconnected from the main power source. Cleaning with the system turned on may be hazardous to the operator and/or destructive to the system.

Exterior Cleaning

The outer surface of the system may be wiped clean with a soft cotton cloth dipped in a mild soap and water solution.

Interior Inspection

Any interior inspection should be performed only by Dune Medical authorized technical personnel.



Warning

- The MARGINPROBE[®] console generates hazardous voltages within the main console.
 - The interior of the system may be serviced only by Dune Medical authorized technical personnel.

Service Information

When communicating with Dune Medical representatives regarding the system, always include the system model and serial number indicated

on the console identification label located on the console's rear panel.



Note,

Warning

Unauthorized servicing or modification of this system may expose the operator or patient to potential high voltage and radiofrequency hazards.

Note

Improper use or maintenance of this system may invalidate the service warranty agreement.

Questions or problems should be referred to your Dune Medical representative, or to the Service Center at:

e-mail: service@dunemedical.com

Caution: Federal law restricts this device to sale by or on the order of a physician

Storage

Do not store the MARGINPROBE[®] system or the sterile probes where they may be exposed to heat, direct sunlight, water or any other liquids. Ensure the console wheels are locked after positioning it in storage.

MARGINPROBE [®] Console		
Temperature	10°C to 40°C / 50°F to 104°F	
Humidity	20% to 85% RH at temperature < 40° C / 104° F	
Altitude	0 to 4572m (0 to 15000 feet)	
Vibration	0.5G maximum, 5Hz to 500Hz	
MARGINPROBE [®] Probe		
Temperature	< 40°C / 104°F	
Humidity	20% to 90% at temperature < 40° C / 104° F	

Caution: Federal law restricts this device to sale by or on the order of a physician

7. TROUBLESHOOTING

Introduction

The MARGINPROBE[®] system is equipped with self-testing software routines that continuously monitor the systems operation. If a system malfunction is detected, an error message will appear on the screen.

The following troubleshooting guides do not attempt to list all possible system failures. Any fault not listed should be referred to Dune Medical Service personnel.



Do not attempt to open or disassemble the system covers.

Troubleshooting Guides

Table 7-1 provides a list of error messages that may appear on the screen, their possible causes and corrective actions to be performed. If the corrective actions listed in the table do not solve the problem, contact your Dune Medical Service representative.

Table 7-1: Error Message	Troubleshooting Guide
--------------------------	-----------------------

Error Message	Probable Cause	Corrective Action
Temperature failure System will shutdown automatically Please contact service	System will shutdown utomatically Please contact service System's internal temperature has risen above 45°C/113°F.	
Fan failure System will shutdown automatically Please contact serviceThe system has detected a fatal failure in the heat dissipation fans.		Contact Dune Medical Service.
Fan problem detected Please notify service at end of procedure	The system has detected a failure in the heat dissipation fans.	Continue normal operation and notify Dune Medical Service.
System Reached Max Number of Measurements. Clear Screen.The screen has displayed its maximal number of measurements.		Clear the screen (long press on T button).

<i>Tuble /-1. Litter message troubleshooling Guide</i> (continued)
--

Error Message Probable Cause		Corrective Action
Please shut system down and Restart	The system has detected an internal error.	Shutdown and restart the system. Resume normal operation. If problem persists, contact Dune Medical Service.
Please Contact Service	The system has detected a fatal internal error.	Contact Dune Medical Service.
Measurement initialization failure Replace Probe	The system detected an error in the probe calibration or measurement process.	Replace the probe and contact Dune Medical Service
Probe failureThe probe data wasReplace Probecorrupted.		Replace the probe and contact Dune Medical Service
Unrecognized Probe was connected	A general problem was found in the probe.	Replace the probe and contact Dune Medical Service
Probe has expired	The probe expiration date has passed.	Replace the probe
Probe has reached time limit #1	The probe has reached its usage limit	Probe can no longer be used for reliable measurements. Replace the probe.
Probe has reached time limit #2	The probe has reached its usage limit	Probe can no longer be used for reliable measurements. Replace the probe.
Probe has reached time limit #3	The probe has reached its usage limit	Probe can no longer be used for reliable measurements. Replace the probe.
Probe has reached its capacity limitThe probe has reached its usage limit		Probe can no longer be used for reliable measurements. Replace the probe.
Probe is not valid with this ConsoleProbe was connected to a different Console in the past		Replace the probe

Table 7-2 lists some possible symptoms of system malfunctions for which no messages are displayed. If the corrective actions listed in the tables do not solve the problem, contact your Dune Medical Service representative.

Symptom	Probable Cause	Corrective Action
Un-requested and/or repetitive measurements when probe is not coupled to tissue.	Vacuum system openings at tip of probe are blocked.	 Try to clear any visible tissue remnants that block the openings at the tip of the probe. If problem persists, replace probe and contact Dune Medical.
Measurement failure Note: If eight consecutive failed measurements are obtained, a message instructing the user to replace the probe will appear on the screen.	 Probe is lifted from tissue before beep is heard. Inadequate coupling of probe to tissue. Coupling obstacle: wires, clips or sutures present on specimen surface at point of measurement. Vacuum airway is disconnected or damaged 	 Leave probe coupled to tissue until beep is heard. Ensure adequate coupling with proper suction by firmly holding probe tip perpendicular to tissue surface. Assure that the probe tip is not placed over wires, clips or sutures. Follow screen instruction and replace the probe if required. If problem persists, contact Dune Medical.
Calibration failure	Probe sensor plate may be covered or obstructed.	 Hold the probe in the air and make sure the sensor is not covered. Click the probe control button to restart calibration Follow screen instruction and replace the probe if required.
The initialization process does not start when turning on the console (a progress bar does not appear on the screen)	Electronic malfunction	 Restart the system. If problem persists, contact Dune Medical.

Symptom	Probable Cause	Corrective Action
Initialization process takes longer than usual (progress bar is shown)	Internal modules are initializing	 Wait 5 minutes to complete initialization. If problem persists, restart the system. If problem persists after restarting the system contact Dune Medical.
Measurement is not triggered upon coupling of probe to tissue	Vacuum airway is disconnected or damaged	 Disconnect and reconnect the probe. If problem persists, restart the system If problem persists after restarting the system, replace the probe. If problem persists after replacing the probe contact Dune Medical.
System is not ready for measurement after connecting the probe Connect Probe message persists after probe connection	 Improper probe connection Probe malfunction. System malfunction. 	 Disconnect and reconnect the probe. If problem persists, restart the system If problem persists after restarting the system, replace the probe. If problem persists after replacing the probe contact Dune Medical.
No sound is heard upon completion of measurement Note: lack of sound does not indicate system malfunction and the system may be used without the audible indication. Screen and LED indications are accurate even in the absence of the audible indication.	 System is muted System speaker malfunction 	 Press volume up button on control panel. (pressing the volume up/down button should produce an audible beep) Continue normal operation using screen and LED indications. Restart system at the end of the procedure and check the sound by pressing the up/down volume button. If problem persists, contact Dune Medical.

Table 7-2: System Malfunction Troubleshooting Guide (Continued)

8. System Specifications

System Performance

• Effective measurement area:	Ø 7.0 mm
• Detection depth:	< 1 mm
• Tissue type:	Detection of in-situ and invasive ductal and lobular breast carcinoma
• System warm-up time:	< 3 minutes
• Measurement time (from tissue attachment to results display):	about 3 seconds

Environmental Requirements

•	Room temperature:	15°C to 30°C / 59°F to 86°F
•	Relative humidity:	30% to 80% at temperature $< 29^{\circ}$ C / 84°F

Physical Specifications

•	Console dimensions [WxDxH]:	60 x 60 x 145 cm 23.6 x 23.6 x 57 inches
•	Console weight:	< 85 Kg. / 187 Lbs.
•	Probe dimensions [LxW]:	197 x 23 mm 7.8 x 0.9 inches
•	Probe box dimensions [WxDxH]:	34 x 25 x 5 cm 13.4 x 9.9 x 2 inches

Electrical Specifications

•	AC Power requirements:	230VAC/50Hz or 115VAC/60Hz, single phase
•	Max AC power:	200 VA
•	Max output to probe:	1 dbm (1mW)

This page is intentionally left blank

Caution: Federal law restricts this device to sale by or on the order of a physician