

Rev. 03

March 2016

# SAFETY AND STANDARDS

350007100

#### MyLab - SAFETY AND STANDARDS

# Introduction

This manual provides information on Safety and Standards for the MyLab product line. This manual consists of the following chapters:

• Chapter 1: Operator's Safety

This chapter describes the situations that could affect the operator's safety when an ultrasound system is used.

• Chapter 2: Patient's Safety

This chapter describes the situations that could affect the patient's safety when an ultrasound system is used.

• Chapter 3: Standards

This chapter lists the standards **MyLab** complies with. It also lists the standards with which the peripherals connected to the device have to comply.

• Chapter 4: Device Labels

This chapter lists the labels provided on the device.

WARNING

#### In this manual a WARNING pertains to possible injuries to the patient and/ or the operator.

CAUTION

# A CAUTION describes the precautions which are necessary to protect the equipment.

Be sure that you understand and observe each of the cautions and warnings.

Keep the manual with equipment for future reference.

#### MyLab - SAFETY AND STANDARDS

# **Table of Contents**

1	Operator's Safety1-1
	Installation Requirements1-1
	Warnings
	Electrical Safety1-1
	Warning1-1
	Caution1-1
	Environmental Safety
	Information about Reusing/Recycling1-2
	Exam Waste1-2
	Moving the Equipment1-2
	Explosive Hazard1-4
	Probes
	Warning1-4
	Caution1-4
	Biocompatibility and Infection Control1-5
	Repetitive Strain Injury1-5
	Working with Video Display1-5
	Wireless Requirements1-6
	Device Modifications1-6
2	Patient's Safety
	Electrical Safety
	Warning
	Electromagnetic Compatibility
	Warning
	Electro-Surgical Devices (ESUs)2-3
	Biocompatibility and Infection Control2-3
	Items in Contact with Patient2-3
	Latex Sensitive Patient
	Probes Superficial Temperature
	Ultrasound Safety
	Introduction
	Clinical Safety2-4
	Ultrasound Bioeffects
	On-screen Real-Time Acoustic Output Display2-5
	The Mechanical Index2-6
	The Thermal Index2-7
	Acoustic Output Display2-7
	The Output Display2-8
	The Output Default Settings2-8
	Methodology and Accuracy of Display2-8
	Maximum Acoustic Output2-9

Davicas Standards	₹_1
Symbols Used in Indices Equations	2-17
-	
Indices Equations	2-16
Definition of Terms	2-14
"In Situ" Intensities Calculations	2-14
Glossary and Definition of Terms	2-14
Acoustic Output Tables	
Which Index When	
Implementing ALARA with MyLab	
Controls Which Indirectly Affect the Intensity	
Controls Which Directly Affect the Intensity	
-	
Acoustic Output Controls	2-10

3	Devices Standards	3-1
	Medical Device Directive	3-1
	Medical Electrical Equipment Standard	3-1
	Electromagnetic Compatibility	3-1
	Biocompatibility	3-1
	Acoustic Output	3-2
	Peripherals Standard Requirements	3-2
	Safety	3-2
	Electromagnetic Compatibility	
	Wireless	3-2
Л	Device Labels	1_1
4		
	Equipment Labels	
	Packaging Labels	4-5

# Chapter

# 1 - Operator's Safety

# **Installation Requirements**

The "Getting Started" manual provides detailed instructions to correctly install and connect your specific **MyLab** model. The same manual also contains all information on the recommended peripherals that may be connected to the system.

If help is needed, Esaote personnel will be glad to provide the necessary assistance to install the system.

#### Warnings

An incorrect installation of the system may cause hazards for the operator. Carefully follow the instructions given in MyLab "Getting Started" manual to install the device.

# **Electrical Safety**

The equipment label, placed on the rear panel, specifies the device electrical requirements. Incorrect connections to the main power may compromise the electrical safety of the system.

#### Warning

Observe the following warnings for maximum safety.

WARNING

 $\square$  GS

- Electrical shock hazard. Do not remove the system or the monitor cover. Refer servicing and internal adjustments to qualified Esaote personnel only.
- Always turn the equipment off before cleaning it.

#### Caution

• To prevent further damage to your system and the accessories, turn the power off if the system does not start up correctly.

AUTION CAUTION Observe these precautions to prevent damage to

your system.

If your system incorporates an LCD and/or a touchscreen, remember that the screen is fragile and must be treated accordingly.

# **Environmental Safety**

#### Information about Reusing/Recycling

This symbol identifies a recyclable component. Depending on the size of the recyclable component, Esaote prints on it this symbol and the indication of the material it is made of.

In this system, packing materials are reusable and recyclable; the casings of the system and the monitor (plastics) and most of the cart components (plastics) are also recyclable.

GS Refer to the MyLab "Getting Started" manual for any additional information on special waste that has to be disposed of according to local regulations.

Exam Waste

Consider any exam waste as potentially infectious and dispose of it accordingly

# Moving the Equipment

G GS MyLab systems are designed to be easily moved by the operator. However, due to the equipment weight, assistance could be needed during transportation. The MyLab "Getting Started" manual details the weight and dimensions of each configuration.

MyLab products can be classified as portable and mobile:

- **Portable** means that the system is equipped with a handle, whose size and weight allow it to be used to carry the system. The term "portable" is always used with this meaning in these manuals.
- A **mobile model** or configuration is equipped with wheels allowing to carry the system from one room to another. The term "mobile" is always used with this meaning in these manuals.

Portable The console can be carried directly by its handle; observe the following precautions:





	• make sure the console is turned off,
	• if built-in, make sure the system screen is secured before and during transportation,
	• disconnect any cable or item (probes, ECG cable) attached to the system,
	• should the console need to be put on the ground, lay it flat,
	• secure the system in a flat position, if transporting it in a vehicle.
Mobile Configuration	The MyLab system complies with EN60601-1: it is not unbalanced by a $10^{\circ}$ inclination. Observe the following precautions when transporting the system:
	• make sure the system is turned off,
	• unlock the cart's wheels before moving the system,
	• avoid unnecessary shocks to the system when rolling it over door jambs or in and out of elevators,
	• when transporting the system with the probes attached, make sure the cables are not dragging on the floor and that the probes are properly positioned in the cart's probe holder,
	• always use the handle to move the system. Never push the system from its sides.
Transportation in Vehicle	Observe the following precautions when transporting the system in a vehicle:
Veniele	• disconnect any cable or item (for example probes, ECG cable) attached to the system and place the probes in their cases,
	• a portable model should be packed in the original shipment case (or other protective devices as available through Esaote) during transportation,
	• for mobile systems, make sure the cart wheels are blocked and the cart secured during transportation.

SAFETY AND STANDARDS

# **Explosive Hazard**

# WARNING The equipment is not suitable for use in the presence of a flammable anaesthetic mixture with air, oxygen or nitrous oxide. Do not use the system in the presence of flammable anaesthetics. Explosion is a hazard under such conditions.

#### Probes

**GS** and **AO** Use only Esaote approved probes with the equipment. The MyLab "Getting Started" manual lists the probes which can be connected to the system. The MyLab "Advanced Operations" manual explains system related special features, when applicable.

The "Probes and Consumables" manual treats all aspects concerning probe cleaning and disinfecting.

#### WARNING

PC

 $\square$ 

Damage caused by dropping a probe, striking it against another object, pinching, kinking or twisting the cable are not covered by the warranty.

# Warning

- If a probe is dropped or suffered an impact against another object, do not use it until an electrical leakage current measurement test has demonstrated that the electrical safety has not been compromised.
- Do not immerse the entire probe in liquid to clean it. The probe is not watertight and immersion may compromise the electrical safety features of the probe.

#### Caution

#### CAUTION

Observe these precautions to prevent damage to your system.

- Never expose the probes to gas, heat or liquid sterilization procedures. These methods can permanently damage the probe.
- Do not connect or disconnect an active probe during live scanning; the system must be in freeze mode or turned off to connect or disconnect a probe.
- Carefully follow the "Probes and Consumables" manual instructions to clean or disinfect a probe.

# **Biocompatibility and Infection Control**

**PC** 

Probes and electrodes intended to be used on intact skin have very limited probabilities to propagate infections; basic procedures as described in the "Probes and Consumables" manual are sufficient for infection control.

Endocavity and transesophageal probes require specific cleaning and disinfecting procedures. See the "Probes and Consumables" manual for complete details on these procedures.

# **Repetitive Strain Injury**

Musculoskeletal disorders have been reported by the clinical literature<sup>1</sup> as a result of repetitive scanning. These musculoskeletal disorders are also referred to as Repetitive Strain Injury (RSI). To prevent the risk of RSI, it has been recommended:

- to maintain a balanced position while scanning,
- not to grip the probe with excessive force,
- to take work breaks to allow muscles to relax,
- to introduce routine exercises such as gentle passive stretching.

# Working with Video Display

Scanning can require long sessions in front of a screen. Consequently visual problems such as eyestrain and irritation can result<sup>2</sup>. Visual discomfort is reduced when the following recommendations are observed:

- orientate the display so that it can be comfortably observed while scanning,
- take rest breaks after a long scanning session.

Necas M. "Musculoskeletal symptomatology and Ripetitive Strain Injuries in Diagnostic Medical Sonographers", Journal of Diagnostic Medical Sonography 12, p. 266-273, 1996
 Pike I, Russo A., Berkowitz J et al. "the prevalence of musculoskeletal disorders

among Diagnostic Medical Sonographers", Journal of Diagnostic Medical Sonography 13, p. 219-227, 1997

See for example OSHA 3092 "Working safely with video terminals display" 1997

# Wireless Requirements

The presence of the following label indicates that MyLab is equipped with built-in wireless capability:



#### This device contains: FCC ID PPD-AR5BHB116 IC 4104A-AR5BHB116

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
- this device must accept any interference received, including interference that may cause undesired operation.



This device contains: FCC ID PD9633ANH IC 1000M-633ANH

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
- this device must accept any interference received, including interference that may cause undesired operation.

WARNING

This device generates and radiates radio-frequency energy. A minimum body to antenna distance of 20 cm must be maintained when the device is installed and operated. The antennas are located on the MyLab systems where indicated by the red circle the next figures.

#### MyLab - SAFETY AND STANDARDS



MyLab Alpha







MyLab Six and MyLabSix CrystaLine



# **Device Modifications**

Esaote is not responsible for any unauthorized modification of equipment (including cables) and/or probes.

WARNING

Do not modify any Esaote equipment without authorization. Always refer to Esaote personnel for authorized modifications of the device.

If the equipment has been modified, appropriate inspections and testing must be carried out to ensure the continued safe use of the equipment.

# Chapter

# 2 - Patient's Safety

# **Electrical Safety**

Warning

Observe the following warning for maximum safety.

WARNING

GS 

- The system must be properly grounded to prevent shock hazards. Protection is provided by grounding the chassis with a three-wire cable and plug; the system must also be powered through a properly grounded receptacle.
- Do not replace the system fuses with different types from those specified by the MyLab "Getting Started" manual.
- Mobile configurations provide insulated plugs and connectors to manage optional hard copy devices (VTR, printers). Follow the instructions in the "Getting Started" manual to install such a device. Incorrect connections may compromise the electrical safety of the system.
- If the operator plans to use hard-copy devices with a portable model, read and carefully follow the instructions in the "Getting Started" manual to install such devices. Incorrect connections or use of peripherals with improper safety characteristics may compromise the electrical safety of the system.
- MyLab models are not watertight and provide a class IP(X)0 degree of protection to liquids; do not expose the system to rain or moisture. Avoid placing liquid containers on the system.
- Remove probes and electrocardiography leads from patient contact before applying a high voltage defibrillation pulse.
- MyLab systems use high frequency signals. Pacemakers could interfere with these signals. The user should be aware of this minimal potential hazard and immediately turn the system

off if interference with the pacemaker operation is noted or suspected.

While using the system in combination with high frequency devices (like electro-surgical devices), be aware that a failure in the surgical device or a damage to the transducer lens can cause electro-surgical currents that can burn the patient. Thoroughly check the system and the probe before applying HF surgical currents to the patient. Disconnect the probe when not imaging.

# **Electromagnetic Compatibility**

Ultrasound systems require special precautions regarding EMC and must be installed and put into service according to the provided information.

Sensitivity to interference is Ultrasound systems are designed to generate and receive radiofrequency (RF) more noticeable in Doppler energy and are, therefore, susceptible to other RF sources. As an example, other medical devices, information technology products or TV/radio transmitters may cause interference with the ultrasound system.

> In the presence of RF interference, the physician must evaluate the image degradation and its diagnostic impact.

Warning

WARNING

modes.

G GS

The "Getting Started" manual provides the table with equipment distance requirements.

- Portable and mobile RF communication equipment may • cause interference with the ultrasound system. Do not use these devices in proximity of ultrasound equipment.
- Use of accessories and cables other than those specified in • the MyLab "Getting Started" manual may result in increased emission or decreased immunity of the system.

If an ultrasound system causes interferences (which can be identified by turning the system off and on) with other devices, the user could try to solve the problem by:

- relocating the system,
- increasing the distance from other devices,
- powering the ultrasound system from an outlet different from the one of the interfering device,

• contacting Esaote Service personnel for help.

#### Electro-Surgical Devices (ESUs)

Electro-surgical devices or other devices that introduce radiofrequency electromagnetic fields or currents into the patient may interfere with the ultrasound image. An electro-surgical device in use during ultrasound imaging will grossly affect the 2D image and render Doppler modalities useless.

# **Biocompatibility and Infection Control**

PC PC

Before each exam properly clean the probes. Refer to the "Probes and Consumables" manual for further details on cleaning and disinfecting probes, kits and electrodes.

#### **Items in Contact with Patient**

Esaote probes and electrodes materials that are in contact with the patient have been proved to comply with EN ISO 10993 "Biocompatibility Tests Requirements", according to their intended use. No negative reactions to these materials have been reported.

#### Latex Sensitive Patient

The USA Food and Drug Administration (FDA) has issued an alert on products composed of latex, because of reports of severe allergic reactions.

#### Note

Esaote probes do NOT contain latex.

WARNING

The probe protective covers used during the patient exam are usually made of latex. Carefully read the protective cover package labeling to check the material used. Be certain to identify latex sensitive patients before starting the exam. Serious allergic reactions to latex have been reported and the user should be ready to react accordingly (for further information refer to the FDA Medical Alert, March 29, 1991, "Allergic Reactions to Latex-Containing Medical Devices"). For additional information in the U.S.A., refer to FDA Medical Alert MDA91-1.

#### **Probes Superficial Temperature**

MyLab has been designed to keep the probes superficial temperature within the limits defined by the IEC 60601-2-37 standard (43°C). Esaote recommends to freeze the system at the end of the exam by pressing the FREEZE key to avoid any

probe overheating. The system will automatically be frozen if left inactive for a few minutes.

The "System Data" section, enclosed in the operator's manual disk, reports the maximum temperature on the probe surface. Before starting the examination, read the "Maximum Probe Temperature" table to identify the probes whose maximum surface temperature can exceed 41 °C.

# **Ultrasound Safety**

#### Introduction

GS GS

MyLab "Getting Started" manual provides data about the acoustic power. Esaote has adopted the more recent requirements and recommendations established by the USA Food and Drug Administration and by the American Institute of Medicine and Biology. MyLab is equipped with the **Acoustic Output Display** feature to provide the user with real time, on-line information on the actual power of the system. The following sections describe the rationale of this methodology. Esaote recommends the use of the **ALARA** principle (see below), which is extensively covered in this manual.

#### **Clinical Safety**

In the USA, in more than three decades of use, there has been no report of injury to patients or operators from medical ultrasound equipment.

#### American Institute for Ultrasound in Medicine (AIUM) Statement on Clinical Safety: October 1982, Revised March 1983, October 1983 and March 1997.

Diagnostic ultrasound has been in use for over 25 years. Given its known benefits and recognized efficacy for medical diagnosis, including use during human pregnancy, the American Institute of Ultrasound in Medicine herein addresses the clinical safety of such use:

No confirmed biological effects on patients or instrument operators caused by exposure at intensities typical of present diagnostic ultrasound instruments have been reported. Although the possibility exists that such biological effects may be identified in the future, current data indicate that the benefits to patients deriving from the prudent use of diagnostic ultrasound outweigh the risks, if any, that may be present.

The **ALARA** (As Low As Reasonably Achievable) principle is the guideline for prudent use: during an exam, the user should use for the shortest duration the least

Refer to the glossary at the end of this chapter for specific terms. amount of acoustic output to obtain the necessary clinical information for diagnostic purposes.

**Ultrasound Bioeffects** 

Although diagnostic ultrasound has an excellent history of safety, it has been known for a long time that ultrasound, at certain levels, can alter biological systems. The AIUM Bioeffects Committee describes two fundamental mechanisms by which ultrasound may induce biological effects: non-thermal or mechanical mechanisms<sup>1</sup> and thermal effects.

Non-thermal bioeffects, also referred to as **mechanical bioeffects**, seem to be caused by the tissue alternate expansion and contraction induced when ultrasound pressure waves pass through or near gas. The majority of these non-thermal interactions, also known as cavitation, deal with the generation, growth, vibration, and possible collapse of microbubbles within the tissue. The occurrence of cavitation depends on a number of factors, such as the ultrasonic pressure and frequency, the ultrasonic field (focused or unfocused, pulsed or continuous), the nature and state of the tissue and boundaries. Mechanical bioeffects are a threshold phenomenon, occurring only when a certain level of output is exceeded. However, the threshold level varies depending on the tissue. The potential for mechanical effects is thought to increase as peak rarefactional pressure increases, but to decrease as the ultrasound frequency increases.

Although there have been no adverse mechanical bioeffects in humans from diagnostic ultrasound exposure, it is not possible to specify thresholds at which cavitation will occur in mammals.

THERMAL BIOEFFECTS

Rise in temperature of tissue exposed to acoustic energy. **Thermal bioeffect** is the rise in temperature of tissue when exposed to acoustic energy. The acoustic energy is absorbed by body tissue; absorption is the conversion of this energy into heat. If the rate of energy deposition in a particular region exceeds the ability to dissipate the heat, the local temperature will rise. The rise in temperature will depend on the amount of energy, the volume of exposure, and the thermal characteristics of the tissue.

**On-screen Real-Time Acoustic Output Display** 

Until recently, application-specific output limits<sup>2</sup> established by the USA Food and Drug Administration (FDA) and the user's knowledge of equipment controls and

"Cavitation" phenomenon

MECHANICAL

BIOEFFECTS

SAFETY AND STANDARDS

<sup>1.</sup> American Institute of Ultrasound in Medicine Bioeffects Comittee "Bioeffects Considerations for the Safety of Diagnostic Ultrasound"; *J:Ultrasound Med*, 1988, 7 Suppl.

Also known as the pre-amendments limits, those values were established on the basis of acoustic output of equipment on the market before 1976.

patient body characteristics have been the means of minimizing exposure. Now, more information is available through a new feature, named the Acoustic Output Display. The output display provides users with information that can be specifically applied to ALARA. It eliminates some of the guesswork and provides both an indication of what may actually be happening within the patient (i.e. the potential for bioeffects), and what occurs when system control settings are changed. This makes it possible for the user to get the best image possible while following the ALARA principle and thus to maximize the benefits/risks ratio.

**MyLab** incorporates a real-time acoustic output display according to the AIUM<sup>1</sup>/ NEMA<sup>2</sup> "Standard for Real-Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment" publication, adopted in 1992 by both institutions. This **output display standard** is intended to provide on-screen display of these two indices, which are related to ultrasound thermal and cavitation mechanisms, to assist the user in making informed risk (i.e. patient exposure)/benefit (diagnostically useful information) decisions. Considering the type of exam, patient conditions and the case study level of difficulty, the system operator decides how much acoustic output to apply for obtaining diagnostically useful information for the patient; the thermal and mechanical indices real-time display is intended to provide information to the system operator throughout the examination so that exposure of the patient to ultrasound can be reasonably minimized while maximizing diagnostic information.

For systems with an output display, the FDA currently regulates only the maximum output. **MyLab** system has been designed to automatically default the proper range of intensity levels for a particular application. However, within the limits, the user may override the application specific limits, if clinically required. The user is responsible for being aware of the output level that is being used. The **MyLab** real time output display provides the user with relative information about the intensity level.

#### The Mechanical Index

The Mechanical Index (**MI**) is defined as the peak rarefactional pressure in MPa (derated by a tissue attenuation coefficient of 0.3dB/cm/MHz) divided by the square root of the probe central frequency in MHz.

With the MI, the user can keep the potential for mechanical bioeffects as low as reasonably achievable while obtaining diagnostically adequate images. The higher the index, the larger the potential. However, there is not a level to indicate that

ODS

Thermal and Mechanical Indices display to assist in making informed risk/benefit decisions.

### MI

Estimates mechanical bioeffects.

<sup>1.</sup> American Institute for Ultrasound in Medicine.

<sup>2.</sup> National Electric Manufacturers Association.

bioeffect is actually occurring: the index is not intended to give an "alarm" but to use it to implement the ALARA principle.

The Thermal Index

TI Relates to temperature rise. The purpose of the Thermal Index (TI) is to keep the user aware of conditions that may lead to a temperature rise under certain defined assumptions. It is the ratio between the total acoustic power to the power required to raise tissue temperature by 1°C, estimated on thermal models. There are currently three thermal indices (each based on a specific thermal model) used to estimate temperature rise whether at the surface, within the tissues, or at the point where the ultrasound is focusing on bone:

- 1. The Soft Tissue Thermal Index (**TIS**) provides information on temperature increase within soft homogeneous tissue.
- 2. The Bone Thermal Index (**TIB**) provides information on temperature increase of bone at or near the focus after the beam has passed through soft tissue.

As with the Mechanical Index, the thermal indices are relative indicator of temperature rise: a higher value represents a higher temperature rise; they indicate that the possibility for an increase in temperature exists and they provide a relative magnitude that can be used to implement ALARA.

#### **Acoustic Output Display**

The acoustic output indices are displayed during live scanning to the right of the screen, together with the transmit power setting.

Indices are displayed in 01.1 increments.

The following abbreviations are used:

Index	Abbreviation
Soft Tissue Thermal Index	TIS
Bone Thermal Index	TIB
Mechanical Index	MI

The output display is organized to provide meaningful information to implement ALARA without "distracting" the user with unnecessary data. During the entry of the patient ID, the user is provided with a choice of applications (Cardio, Abdominal, Reproduction, etc.); depending on the selection, the system will default the appropriate indices.

#### Note

Index values below 0.4 are NOT displayed by this system.

To optimize ALARA, index values equal or higher than 0.4 are displayed even if the maximal index value does not exceed 1.0.

The Output Display

The following table shows the indices used for each clinical application. Indices are displayed in 0.1 increments.

Application	MI	TIS	TIB
OB (Reproduction)	Yes	Yes	Yes
Endorectal	Yes	No	No
All others	Yes	Yes	Yes <sup>a</sup>

a. Only when TIB≠TIS

The Output Default Settings

System default settings depend upon the probe, the mode of operation and the application which is selected during the patient ID procedure. The MyLab defaults the transmit power to obtain output levels that are below the historic Ispta limits established by the FDA for the selected application.

#### Methodology and Accuracy of Display

The displayed indices values must be interpreted as relative information to help the user to achieve the ALARA principle.

Initial data are derived from laboratory measurements based on the AIUM standard. Then the indices are calculated beginning from these measurements according to the AIUM/NEMA "Standard for Real-Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment" publication. Many of the assumptions used for measurements and calculation are conservative in nature. The measured water tank values are derated using the conservative attenuation coefficient established by the standard (0.3 dB/cm/MHz). Over-estimation of actual in-situ exposures is thus part of the calculation process.

In combined modes (ex.: 2D+Doppler), the indices will show the highest value between the two modes.

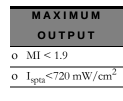


- o Accuracy: ±14% for the MI
- o  $\pm 30\%$  for the TI

A number of factors influence the estimation of the accuracy of the displayed indices, the most significant ones being the variability between probes and the laboratory measurements accuracy (hydrophone, operator, algorithms, etc.) itself, while variability of the system pulsar and efficiency is a minor contributor.

The accuracy estimate, based on the variability range of probes and systems, and on the inherent modeling and measurements errors, is 14% for the MI and 30% for TI indices; this accuracy estimate does not consider errors in/or caused by measuring with the AIUM standard.

Maximum Acoustic Output



This system does not use the historic FDA limits for Isppa and Imax, but rather the MI, which is now considered a better relative indicator of non-thermal bioeffect mechanisms. The maximum MI is below 1.9; the FDA has recognized this value as equivalent to pre-amendments Isppa limits. The maximum output for Ispta is limited to the pre-amendments FDA limit for peripheral vascular applications (720 mW/cm<sup>2</sup>).

Application	Preamendments Ispta Limits (mW/cm2)	MyLab Maximum (mW/cm2)
OB/Fetal (Reproduction)	94	430
Cardiac	430	720
Other	94	720

Other application limits have been established as per this table:

The maximum output for a given probe can be less than the system limit, since the maximum depends on various elements (for example crystal efficiency, mode of operation).

**Acoustic Output Controls** 

Control features may be divided into three categories:

- 1. controls which directly affect the intensity (direct controls),
- 2. controls which indirectly affect the intensity (indirect controls),
- 3. controls, which do not affect the intensity, such as the gains and the processing curves.

#### **Controls Which Directly Affect the Intensity**

This category includes two system controls:

#### DIRECT CONTROLS

- o the application
- o the power

- the application selection, which establishes the appropriate range of intensities (see maximum output section); the application also establishes the indices to be displayed;
- the **POWER** control, which allows an increase or decrease in the output intensity within the range of the selected application. This parameter will affect both the MI and the TI values.

#### **Controls Which Indirectly Affect the Intensity**

This category includes controls, which change several aspects of the transmitted ultrasonic field rather than the intensity. Intensity is affected because of the field variations. Each mode has its own pulse repetition frequency (PRF) and intensity level; moreover, for each mode, a number of parameters will indirectly affect the transmitted field.

#### Note

The TI index display depends on the application and on the mode.

The MI may increase whenever the PRF is decreased, i.e. when the field of view is increased.

**MyLab** allows the user to set the transmit focal point which will affect both indices by varying the beam profile. Generally, higher MI's and TI's will occur with closer focal points. If more than one transmit focal point is activated, MI and TI values will each correspond to the zone with the largest value. In addition, all system probes can image at two frequencies; both indices are usually different, depending on the probe bandwidth.

Tissue<br/>Enhancement<br/>Image (TEI)The same controls described for 2D affect the acoustic output. Because the tissue<br/>response is a non-linear phenomenon, this modality usually requires higher<br/>acoustic outputs than conventional imaging. While using this mode, the MI is<br/>your primary concern; a deeper transmit focal point helps to keep the MI value as<br/>low as possible.

M-Mode In M-Mode, the transmitted field is only affected by the transmit focal point and the frequency. If M-Mode is displayed with 2D and the 2D is updated, the system may show the latter mode MI (and TI if available) if higher.

# 

INDIRECT

- o PRF
- o Focal Point
- o Frequency
- o CFM Process
- o Sample Volume

2D

Color Flow Mapping (CFM)	The MI is primarily dependent on 2D settings, i.e. the depth (which will determine the 2D and color PRF) and the transmit focal point. The MI may also be increased by a decrease in the color PRF. The TI may be increased by increasing the color CFM. Increasing the color frame
	rate may increase the TI while decreasing the MI. Finally, probes can provide color at two frequencies; the outcome in terms of transmitted field is marginal and largely unpredictable.
Tissue Velocity Mapping (TVM)	This mode optimizes CFM settings in order to image the movement of tissue, thus the same controls described for 2D-CFM affect the acoustic outputs.
Pulsed Wave Doppler (PW)	In PW, the sample volume depth automatically sets the Doppler PRF and the focal point. Deeper sample volumes will cause lower PRF; the MI may, however, not increase since the focal point is far, while the TI is generally reduced. The TI may, however, change if the sample volume size is varied. This factor accounts generally for a MI modification.
Tissue Velocity (TV)	The TV Doppler optimizes your settings to analyse tissue motion.
	Finally, most probes provide Doppler at two frequencies; the outcome in terms of transmitted field is marginal and largely unpredictable.
Continuous Wave Doppler (CW)	In CW, the only "variable" factor is the Doppler frequency. As stated before, most probes provide Doppler at two frequencies; the outcome in terms of transmitted field is marginal and largely unpredictable. The user can vary the spectral velocity range; this does NOT, however, change the system's PRF.

#### Note

In Doppler modes, if the tracings are displayed with an updated 2D, the 2D values are used if higher than the Doppler indices.

Implementing ALARA with MyLab

Prudent use implies that during an exam the user should use for the shortest time the least amount of acoustic output to obtain the necessary clinical information for diagnostic purposes. In other words, the goal is to keep the TI and the MI indices as low as possible for the shortest time while obtaining the necessary clinical information. This section does not cover the patient and technique factors, which may influence the indices such as the patient body size, the tissue perfusion characteristics, the presence or the absence of fluid, etc.

#### **ALARA Guidelines**

- Select the appropriate application when you enter the patient data.
- Depending on the patient characteristics and the type of exam, select the appropriate probe and frequency.
- Use the system capabilities to preset the MyLab system to default each mode according to your needs or specific applications; this will reduce the need for real time interactions and help to quickly obtain useful images, thus reducing ultrasound exposure.
- Start scanning with a low output level and optimize the focusing, the gains and all other system adjustments; if this is not adequate for diagnostic purposes, then increase the output level. In cardiac studies, use Tissue Enhancement Imaging if acoustic noise is affecting the images' readability.
- Use the output display feature to guide your settings; remember that the indices do not consider TIME exposure: the higher your indices, the shorter the patient exposure should be.

#### Which Index When

In **cardiac**, general imaging (**abdominal**) and **musculoskeletal** exams, the system displays the TIS in addition to the MI. In imaging and CFM modes, the primary concern is in keeping the "cavitation" predictor as low as possible. You can minimize the MI by reducing the power to the lowest possible level, and adjusting the TGC and general gain controls. Use the transmit focal point to enhance resolution and sensitivity in the area of interest: this may increase the MI, but because of the enhanced sensitivity, you may be able to reduce the transmit power, thus reducing the MI. Decreasing the imaging depth as low as possible may allow the system to increase the PRF and thus reduce the MI.

In Doppler modes, if you are working with a 2D + Doppler display, the MI will show the 2D value (because it is higher than the Doppler one) and the Doppler TIS; the latter parameter should be your primary concern: the MI value reflects the energy to which the patient is exposed only for a minimal time, i.e. between every sweep. You may want however to remember that whenever varying the Doppler speed: increasing the speed will cause the 2D to be refreshed more often. You may

In cardiac and abdominal examinations, MI is the primary concern in imaging modes, while the TIS is the principle index in Doppler.

GS

See the *"Getting"* 

your system.

Started" manual for

eventually freeze the 2D or switch to a full screen mode; however, this will probably increase the time to actually find the desired signal, and therefore the exposure time.

In OB, the TIB should be considered when scanning a second or third trimester fetus, while the TIS is more reliable for earlier exams. In **OB** (**Reproductive**) exams, this system displays both the MI and the TIB in imaging and CFM modes. While the MI will remain your primary concern in those modes, you should also consider the TIB in imaging a second or third trimester fetus as a conservative estimate of the actual temperature rise. In PW Doppler, the latter value is the primary parameter to consider for second or third trimesters pregnancies while the TIS is a more reliable indicator for earlier exams. The general guidelines already expressed for the previous exams remain valid.

#### **Acoustic Output Tables**

According to the IEC61157 and EN 60601-2-37, the acoustic output tables give the acoustic output data for each probe in every operating mode. These tables are in the MyLab Operator Manuals disk.

# **Glossary and Definition of Terms**

"In Situ" Intensities Calculations

The intensity measurements made in water in the laboratory must be derated to reflect the effects of attenuation. When determining the possible effects of the ultrasound beam on tissue, the intensity encountered at the tissue site must be calculated. Because of attenuation of the beam within the body, the intensity at the tissue site ("in situ") may be 10 to 100 times less than if it was measured at the same location in water. The amount of attenuation from experience by an ultrasound beam as it travels through the body tissue is determined by three factors:

- 1. Type of tissue along the beam path,
- 2. Frequency of the ultrasound energy,
- 3. Distance covered by the beam.

In order to achieve a conservative approximation of attenuation due to these three factors, the FDA requires the application of the following formula:

 $Id = Iw \exp \left( -0.23 a f z \right)$ 

• I<sub>d</sub> is the estimated "in situ" intensity at the tissue site,

• $I_w$ is the intensity measured in water at a distance "z", measured in cm,
• a is the attenuation coefficient <sup>1</sup> expressed in dB/cm/MHz,
• f is the acoustic frequency in MHz of the ultrasound beam.
Definition of Terms
The <b>acoustic intensity</b> generated by an ultrasound probe is usually described as follows:
The Spatial Peak Time Average Intensity is an ultrasound intensity averaged over time at the point in the acoustic field where the pulse average intensity is at maximum.
The Spatial Peak Pulse Average Intensity is an ultrasound intensity averaged over the pulse transmission time at a point in the acoustic field where the pulse average intensity is at maximum.
The Maximum Intensity is an average intensity during the half-cycle with the greatest amplitude during the pulse.
The Mechanical Index is defined as the peak rarefactional pressure in MPa (derated by a tissue attenuation coefficient of $0.3 \text{ dB/cm/MHz}$ ) divided by the square root of the probe central frequency in MHz.
The Thermal Index is the ratio between the acoustic power and the power required to raise tissue temperature by 1°C, estimated on thermal models.
The peak rarefactional pressure (pr in MPa) is the temporal peak rarefactional pressure amplitude at a specified point.
The Pulse Intensity Integral (PII) is the time integral of instantaneous velocity for any specific point and for any specific pulse, integrated over the time in which the envelope of acoustic pressure or the envelope of hydrophone signal for the specific pulse is non-zero. It is equal to the energy fluence per pulse.

1. As per the FDA, this coefficient is equal to 0.3dB/cm/MHz

#### **Indices Equations**

Parameter	Equation
Soft Tissue at Surface	•
TIS (scanned <sup>a</sup> )	$\frac{100}{210}$
TIB (scanned <sup>a</sup> )	$\frac{\frac{W_{01}}{210}}{f_c}$
(••••••••• )	- 0
Large Aperture ( $A_{aprt} > 1 \text{ cm}^2$ )	
TIS (unscanned <sup>b</sup> )	$\frac{max_{z>zbp}[min(W_{.3}(z);I_{TA.3}(z) \times 1cm^{2})]}{210}$
	210
	$\frac{210}{f_c}$
Small Aperture (A <sub>aprt</sub> ≤ 1 cm <sup>2</sup> )	
TIS (unscanned <sup>b</sup> )	$\frac{W_0}{210}$
	$\frac{\frac{w_0}{210}}{f_0}$
	J c
Bone at Focus TIB (unscanned <sup>b</sup> )	
	$min\left\{\frac{\sqrt{W_{.3}(Z_{B.3})I_{TA.3}(z_{B.3})}}{50};\frac{W_{3}(z_{B.3})}{4,4}\right\}$
	where $z_{B,3}$ is the depth that maximizes
	$W_{.3}(z)I_{TA.3}(z)$ , or, equivalently, the depth of
	I <sub>SPTAB.3</sub> .
Mechanical Index (MI)	$\frac{I_{SPTAB.3.}}{\frac{p_{r.3}(z_{sp})}{\sqrt{c}}}$
	$\sqrt{f_c}$
	where $p_{r,3}(z_{sp})$ is the peak rarefactional pressure (in
	MPa) derated by 0.3dBcm <sup>-1</sup> MHz <sup>-1</sup> to the point on
	the beam axis $z_{sp}$ where pulse intensity integral
	(PII.3) is maximum, and $f_C$ is the center frequency
	(in MHz).

a. The scanned mode (or autoscanning) is the electronic or mechanical steering of successive ultrasonic pulses or series of pulses, through at least two dimensions.

b. The unscanned mode (or nonautoscanning) is the emission of ultrasonic pulses in a single direction, where scanning in more than one direction would require moving the transducer assembly manually.

Symbol	Definition
$A_{aprt}$ (cm <sup>2</sup> )	Active aperture area
$\frac{A_{aprt} (cm^2)}{d_{eq}(z) (cm)}$	Equivalent beam diameter
1	$\overline{4W_3(z)}$
	$\sqrt{\frac{4W_{.3}(z)}{\pi I_{TA,3}(z)}}$
	1110
$D_{eq}$ (cm)	Equivalent aperture diameter
	$\sqrt{\frac{4A_{aprt}}{\pi}}$
	$\sqrt{-\pi}$
f <sub>c</sub> (MHz)	Center frequency.
$I_{SPTAB.3} (mW/cm^2)$	Equivalent to the spatial peak temporal average
511110.5 \ /	derated (0.6 dBcm <sup>-1</sup> MHz <sup>-1</sup> ) intensity
$I_{TA.3}$ (z) (mW/cm <sup>2</sup> )	Temporal average intensity derated to depth z
$rac{I_{TA.3} (z) (mW/cm^2)}{W_0 (mW)}$	Time average acoustic power at source
W <sub>01</sub> (mW)	Time average acoustic power at the source emitted
	from the central centimeter of the active aperture
W.3 (z) (mW) (mW/cm)	Time average acoustic power derated to depth z
(mW/cm)	Acoustic power per unit linear length (for example
	of a linear array)
z (cm) z <sub>bp</sub> (cm)	Depth from the surface along the beam axis
z <sub>bp</sub> (cm)	Break point depth (minimum depth for intensity
	measurements in the TIS (unscanned) models)
	$z_{bp} = 1.5D_{eq}$
z <sub>B.3</sub> (cm)	Depth of the maximum temperature rise in the bone
	at focus model
p <sub>r.3</sub> (zsp)	Peak rarefactional pressure (in MPa) derated by 0.3
	$dBcm^{-1}MHz^{-1}$ to the point on the beam axis $z_{sp}$
	where pulse intensity integral (PII.3) is maximum

# Symbols Used in Indices Equations

# Chapter 3

# 3 - Devices Standards

# **Medical Device Directive**

This system complies with the Medical Device Directive (MDD) 93/42/EEC and its amendments 2007/47/EEC, according to which Esaote has classified this device as a Class IIa device.

#### Note for U.S. Customers

U.S. Federal Law restricts these devices to sale, distribution and use by or on the order of a physician.

# Medical Electrical Equipment Standard

As defined in EN60601-1 (IEC Standard 60601-1, Safety of Medical Electrical Equipment), **MyLab** models are classified as Class I, with applied parts of type B or BF (probes), and of Type CF (ECG).

These devices also comply with the EN 60601-2-37 (IEC 60601-2-37) "Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment".

# **Electromagnetic Compatibility**

🖾 GS

Each MyLab model complies with the EN60601-1-2 (Electromagnetic Compatibility). Refer to the MyLab "Getting Started" manual for the electromagnetic emissions classification of the devices and electromagnetic immunity compliance levels.

# **Biocompatibility**

The probe and electrode materials that are in contact with patients, comply with the applicable requirements of EN ISO 10993-1, according to their intended use. No negative reactions to these materials have been reported.

# **Acoustic Output**

MyLab global maximum acoustic output doesn't exceed the FDA pre-amendment upper limits (refer to "Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers", issued on September 2008).

# **Peripherals Standard Requirements**

When peripherals are connected to an ultrasound system, they become part of a medical system. Therefore they must comply with the below mentioned standards to maintain the overall system conformity.

Safety

#### The peripheral device must:

meet the EN60601-1 OR in accordance with EN60601-1-1:

- the device must meet the applicable safety standards for its category
- the device must be powered through an isolation transformer designed for medical applications.

If **MyLab** is equipped with the cart equipped with isolation transformer, the isolation requirement is fulfilled by powering the device through one of the cart's insulated plugs.

**Electromagnetic Compatibility** 

The peripheral device must:

- meet the EN55011 or EN55022 emission limits, according to the environment where the system is used and
- meet the EN50082-1 or EN61000-6-1 immunity requirements.

### Wireless

**MyLab** complies with the Radio equipment and Telecommunications Terminal Equipment Directive 1999/5/EC and is CE marked.

MyLab is a device in Class 2 according to R&TTE Directive.

If MyLab is equipped with wireless capability, be informed that:

- whenever the system is used in a hospital, the use of wireless devices might be restricted.
- wireless devices can operate in European countries without restrictions indoor; their use is subjected to restricted frequency band outdoor in France. Refer to local regulations for further information.

WARNING

The use of wireless devices might be restricted in certain locations: always verify local regulations before using it.

#### MyLab - SAFETY AND STANDARDS



# 4 - Device Labels

MyLab devices use the EN60601-1 safety symbols for medical electronic devices to classify a connection or to warn of any potential hazard.

The following tables describe the labels that may be used on the equipment and the packaging.

# **Equipment Labels**

Symbol	Explanation
	On (Mains Power) <sup>a</sup>
0	Off (Mains Power) <sup>b</sup>
ф	Fuse
۲	Type CF applied part
Ŕ	Type B applied part
Ŕ	Type BF applied part

Symbol	Explanation
$\checkmark$	Equipotential connection (it may be found in an optional cart)
4	Dangerous Voltage
$\triangle$	This symbol generically means "Caution". Consult the appropriate sections of user manuals.
	General Warning
Ĺ	Operating instructions. This symbol advises to carefully read the user manuals.
8	Operating instructions. This symbol indicates to carefully read the user manuals.
F©	Device contains radio module fully compliant with CFR47 Part 15 Sub.C (under FCC rules). Device meets requirements of CFR47 Part 18 (under FCC rules).
()	Alert sign. Restrictions on the use of the radio module apparatus may apply in some countries or geographic areas.
	Separate collection for electrical and electronic equipment. This symbol indicates to dispose of the equipment as special waste according to the applicable local regulations.

Symbol	Explanation
IP	<ul> <li>This symbol indicates the degree of protection provided by the enclosure per IEC 60529.</li> <li><b>IPX1</b>: Protected against vertically falling water drops.</li> <li><b>IPX7</b>: Protected against the effects of temporary immersion in water.</li> <li><b>IPX8</b>: Protected against the effects of continuous immersion in water.</li> </ul>
MOD.	Device model
REF	Device part number
SN	Device serial number
~~~	Date of manufacture. The date is located adjacent to the symbol.
	Manufacturer. The date of manufacture can be combined with this symbol.
CE	CE Mark of Conformity
	Product certified by CSA for U.S. and Canadian markets
•	USB connection

Symbol	Explanation
	Headphone connection
Q,	Microphone connection
	ECG connection
$\bigcirc$	Output connection
$\rightarrow$	Input connection
$\bigcirc$	Combined Input/Output connection
	LAN connection
	Direct current
$\sim$	Alternating current

Symbol	Explanation
(((••)))	Radio frequency signal

a. Not available on MyLab Alpha and MyLab Gamma console.

b. Not available on MyLab Alpha and MyLab Gamma console.

# **Packaging Labels**

Symbol	Explanation
X	Temperature limit
) (%)	Humidity limit
<b>(</b>	Atmospheric pressure limitation
Â⊡	Stacking limit
Ţ	Fragile: handle with care
Ť	Keep away from rain

Symbol	Explanation
<u>††</u>	This way up. It indicates the correct upright position.
子	Use no hooks
ES I	Recycling material
	Recycling material. The packaging material complies with the RESY requirements.