

BINDEX

Osteoporosis Diagnostics



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The Quality Management System of Bone Index Finland Ltd. complies with the United States Food and Drug Administration Code of Federal Regulations Title 21 Part 820 (Quality System Regulation), the European Medical Device Regulation 2017/745 (MDR) and the Quality Management Standard EN ISO 13485:2016. Bindex products comply with the Medical Device Directive MDD 93/42/EEC requirements.

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2. Warnings and precautions

Before using Bindex, the user must read and understand the following safetyrelated information. The user shall adhere to the warnings to ensure a safe and reliable performance of the system.



ELECTROMAGNETIC ENVIRONMENT HAZARD:

The Bindex system needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in Appendix: Guidance and manufacturer's declaration -Electromagnetic Compatibility. If the use environment has specific requirements for EMC, the system setup should be assessed and inspected by qualified personnel. Incorrect installation could result in increased electromagnetic emissions or decreased electromagnetic immunity of this device and result in improper operation.



ELECTROMAGNETIC ENVIRONMENT HAZARD:

Portable and mobile radio frequency (RF) communications equipment can affect the Bindex BI-2 device and result in improper operation. Do not use such equipment near the Bindex BI-2 device (See Appendix: Guidance and manufacturer's declaration - Electromagnetic Compatibility for additional information).



ELECTRICAL HAZARD:

Equipment used in the Bindex measuring system must comply with IEC 60601-1:2005+A1/2013 (medical electrical equipment), IEC 62368-1:2014 (non-medical IT equipment) or their general IEC/ISO variants. Electrical safety of the system can only be guaranteed on compliant equipment.



ELECTRICAL HAZARD:

Do not connect or use other devices in the Bindex measurement computer during patient measurements (excl. the mouse, keyboard and display). The electrical and electromagnetic compatibility of the system can only be guaranteed when Bindex device alone is connected.



ELECTRICAL HAZARD:

Non-medical equipment (including the PC) should be located outside the patient environment as described in IEC 60601-1 to guarantee the electrical safety of the system. If it is necessary for the non-IEC 60601-1 compliant equipment to be located within the patient environment, that equipment shall be powered by an internal battery or an EN 62368-1:2014 compliant isolation transformer or be connected to system ground via an additional protective earth terminal.



ELECTROMAGNETIC ENVIRONMENT HAZARD:

WARNING: Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.



HEALTH HAZARD:

Do not make Bindex measurements on the surface of skin with open sores or broken or irritated skin. There is an infection risk.



INCORRECT USE HAZARD:

Do not use the Bindex device in case of a fractured bone at measurement location. The result is only reliable when the measurement is made on an intact bone.

INCORRECT USE HAZARD:

Do not use the Bindex device in case of implants, plates or fixations at measurement location. If possible, conduct the measurement on the other leg. The result is only reliable when the measurement is made on an intact bone.



ENVIRONMENTAL HAZARD:

Do not use the Bindex device outdoors. The device is intended for indoor use only. See 6 Operating environment.

ELECTRICAL HAZARD:

Do not use the Bindex device near a heat source or an air conditioner. This may cause condensation of moisture inside the equipment. Moisture may cause improper operation or electrical safety hazards due to short circuiting. Let the device settle to operating room temperature before operation.



HEALTH HAZARD:

Use only gel intended for clinical ultrasound coupling for measurements with the Bindex device. Improper gel or other substances may cause symptoms such as skin irritation to the patient or operator.



INCORRECT USE HAZARD:

Do not apply ultrasound gel on the surface of the Bindex transducer before calibration. Conducting the calibration with an unclean transducer may cause the calibration or the initial device setup to fail. See 8.6 Bindex quality verification.



INCORRECT USE HAZARD:

Always use the Bindex BI-41 Measure for determination of the proper measurement location. The location is standardized for this measurement to produce reliable results. Failing to use the Bindex Measure in determining the location may lead to incorrect results.



MECHANICAL/ELECTRICAL HAZARD:

If you drop or bump the device on hard surfaces, conduct quality verification measurements. In case of any visible mechanical damage, please contact your local distributor or Bindex Support and Service (see for service. Store the device in its protective case between uses. Do not use a damaged device!



ENVIRONMENTAL/FIRE HAZARD:

The Bindex device is not intended to be used in oxygen rich environment. Using the device in an oxygen rich environment may lead to a fire or explosion.



ELECTRICAL HAZARD:

The patient shall be informed not to touch the connectors of the ME system (e.g. laptop connectors) during measurements. Touching the connectors of the system results in potential electrical safety hazard.

Where possible, the computer should be kept away from the patient area.



ELECTROMAGNETIC ENVIRONMENT HAZARD:

WARNING: Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.



ELECTROMAGNETIC ENVIRONMENT HAZARD:

WARNING: Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the Bindex device, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.



HOME ENVIRONMENT HAZARD: WARNING: Keep away from pets.

HOME ENVIRONMENT HAZARD: WARNING: Keep out of reach of children.



3. Symbols and Abbreviations

3.1. Abbreviations

US	Ultrasound
DI	Density Index
DXA	Dual Energy X-ray Absorptiometry
BMD	Bone Mineral Density
Cth.	Cortical thickness

3.2. Symbols glossary

The symbols of this glossary can be found in Bindex product labels and this User Manual. If unsure about the meaning of a symbol, please contact Bindex Support and Service (see ch. 12 Contact information)

DEVICE SYMBOLS



Refer to instruction manual/booklet. To signify that the instruction manual/booklet must be read. (ISO 7010, Ref.No. M002)



Serial number. Indicates the manufacturer's serial number so that a specific medical device can be identified. (EN ISO 15223-1:2016, Ref.No. 5.1.7)



Manufacturer. Indicates the medical device manufacturer name and address. (EN ISO 15223-1:2016, Ref.No. 5.1.1)



Type BF applied part. To identify a type BF applied part complying with IEC 60601-1. (IEC 60417, Ref.No. 5840)



Caution. Indicates the need for the user to consult the instructions for use for important cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device itself. (EN ISO 15223-1:2016, Ref.No. 5.4.4, symbol variant 0434B)



The device is USB (Universal Serial Bus) compatible. The cable is equipped with a USB Type A connector. (Universal Serial Bus Specification rev.1.1, Fig. 6-5)



Temperature limit (for declared conditions). Indicates the temperature limits to which the medical device can be safely exposed.

(EN ISO 15223-1:2016, Ref.No. 5.3.7)



Consult instructions for use. Indicates the need for the user to consult the instructions for use. Includes the URL (web address) to the electronic User Manual. (EN ISO 15223-1:2016, Ref.No. 5.4.3)



Class II device. To identify equipment meeting the safety requirements specified for Class II equipment according to IEC 61140. Referring to electrical equipment in which protection against electric shock does not rely on basic insulation only, but in which additional safety precautions such as double insulation or reinforced insulation are provided, there being no provision for protective earthing or reliance upon installation conditions. (IEC 60417, Ref.No. 5172)



IP rating. Indicates the protection level of the device against solid foreign objects and water. Bindex is rated as IP22, indicating protection against solid foreign objects of 12,5 mm and greater in diameter (e.g. fingers), and vertically falling water drops when the device is tilted up to 15° (IEC 60529)



Symbol for the United States only, not applicable in the European Union or the United Kingdom. (21 CFR 801.109 (b) (1))

PACKAGE SYMBOLS



EU only: Separate collection for electrical and electronic equipment waste required (see ch. 11.1 Disposal). (EN 50419:2006)



Fragile, handle with care. Indicates a medical device that can be broken or damaged if not handled carefully. (EN ISO 15223-1:2016, Ref.No. 5.3.1)

Keep dry. Indicates a medical device that needs to be protected from moisture. (EN ISO 15223-1:2016, Ref.No. 5.3.4)

In this manual, software controls are indicated as follows: BUTTONS in capital letters, Windows and Pages in bolded capital and small letters and Editable *fields* in italic typeface.

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4. Indications for use

Bindex measures apparent cortical bone thickness at the proximal tibia and can be used for assessment of probability of osteoporosis risk. Results can recommend that the patient consult with a physician for osteoporosis evaluation and diagnosis. When done by a healthcare professional, Bindex measurement can be used in conjunction with other clinical risk factors or patient characteristics as an aid to the physician in the diagnosis of osteoporosis and other medical conditions leading to reduced bone strength and in the determination of fracture risk.

4.1. Intended use

Bindex measures apparent cortical bone thickness at the upper shaft of tibia (See Figure 1) and reports the diagnostic parameter, Density Index (DI), an estimate of hip Bone Mineral Density measured with gold standard Axial DXA. Thresholds for osteoporosis for DI have been determined in comparison to DXA.

After the measurement, Bindex Software gives an estimation of the presence of osteoporosis marked in the color bar: Green (Low Probability of Osteoporosis), Yellow (Additional Investigations Needed) or Red area (High Probability of Osteoporosis). A total of 90% of osteoporotic patients diagnosed by hip BMD are in the yellow or red area (90% sensitivity) and 90% of non-osteoporotic patients are in the green or yellow area (90% specificity). Statistically at least 80% sensitivity and specificity for hip osteoporosis will be reached with 95% confidence. Patient classification is based on thresholds (separating red/yellow/green areas) published in a study by Karjalainen et al. "New method for point-of-care osteoporosis screening and diagnostics" in Osteoporosis International 2016.

A DI value below the upper threshold indicates for consultancy of a physician for osteoporosis evaluation and diagnosis. When done by a healthcare professional, the DI reported by Bindex can also be used as an aid to the physician in osteoporosis diagnostics and estimation of fracture risk.

Currently the use of Bindex DI thresholds is validated for Caucasian and Hispanic

women at the age between 50 to 90 years. Bindex measurement takes about one minute. Bindex device should be operated by a physician, nurse, pharmacist or trained person with a suitable background education and skills.

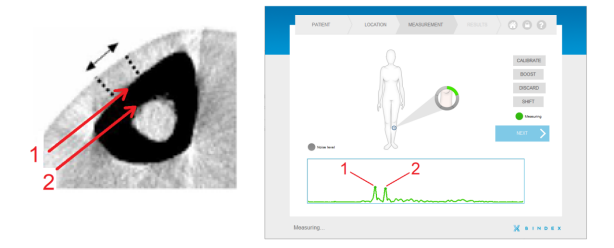


Figure 1: A Computed Tomography image of tibia shows the tubular structure of the bone (black area). When measuring the tibia with Bindex, the ultrasound echoes back from the front (1) and the back (2) surface of cortical bone layer. These echoes need to be clearly distinguishable. Bindex will automatically accept the echoes.



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5. Bindex BI-2 overview and technical specification

Bindex BI-2 device

The Bindex BI-2 device consists of the handpiece including the measurement electronics and the USB cord (Figure 2). The device is connected to a free USB port of a personal computer. In the device, an electrical pulse is sent to the transducer which transforms the pulse into ultrasound waves that are transmitted into the bone. The transducer collects the sound waves reflected from the bone and transmits the signal via the electronics to the PC software for analysis.



Figure 2: Bindex BI-2 device shown together with the BI-41 Measure.

Bindex Software

Bindex utilizes software which is provided to customer on a USB drive or as a downloadable installation package. The software can be installed on Windows 8.1 or Windows 10 operating systems on a PC. The Bindex device is operated using the software GUI (Graphical User Interface) to control the pulser and to collect the measured signals. Signals are analyzed to calculate the diagnostic/screening parameter DI. Results are saved in the Bindex database and can be exported in PDF format or as a text file for easy transfer to e.g. a

spreadsheet program.

Bindex BI-41 Measure

For determination of the standard location for Bindex measurement, device is supplied with a custom stick measure (Figure 3). The Bindex Measure is always used to determine the ultrasound measurement location at 1/3 length of the proximal tibia.

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Figure 3: The Bindex Measure is used for determination of standard measurement location at the tibia.

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Specifications

Мес	chanics
Weight (incl. USB cord)	128g
Size (handpiece)	119 x 42 x 34mm (length x width x height)
USB cord length	2.0 m
IP Rating	IP22
	(Protected against: - Solid foreign objects of 12,5 mm and greater in diameter: - Vertically falling water drops when the device is tilted up to 15°)
Ele	ectrical
Power supply	Powered from PC USB port, 5V
Fuse	6V / 500 mA, resettable (not serviceable by user)
Envir	onmental
Operating Temperature	+15+ <mark>4</mark> 0 °C
Storage Temperature	+15+40 °C
Transport Temperature	-25+60 °C
Atmospheric Pressure	700hPa to 1060hPa (mbar)
Humidity	590%
Ultr	asound
Transducer centre <mark>fr</mark> equency	3.0 MHz
Transducer type	Focused
Mechanical Index	0.220
Thermal Index (TIB _{bs,ns})	0.011
Spatial-peak temporal-average intensity (I _{spta})	6.5 mW/cm ²

Safety standards compliance

Medical electrical equipment safety	EN 60601-1:2006
Electromagnetic compatibility	EN 60601-1-2:2015
Use in Home Healthcare Environment	EN 60601-1-11:2015
Ultrasound safety	EN 60601-2-37:2008, EN 62359:2011

Bindex and the connected PC are together considered a medical electrical system. The computer power source must comply with the IEC 62368-1:2014 standard, otherwise it is mandatory to connect the PC operated with Bindex to the mains supply with a medical isolation transformer. An isolation transformer or an additional protective earth connection from the computer is also required when the computer does not comply with IEC 60601-1:2005 and it is used within the patient environment.

Bindex can also be used with an IEC 62368-1:2014 compliant laptop computer operating on battery power without connecting to the supply mains. In this case, no additional precautions concerning electrical safety are required.



ELECTRICAL HAZARD

The Bindex system, including the computer, must only be used with the electrical setup specified in this chapter. Operator and patient safety regarding electrical hazards can only be guaranteed with the specified conditions and devices.



ELECTRICAL HAZARD:

DO NOT plug the Bindex BI-2 device into supply mains!



6. Operating environment

See section 5 for operating and storing conditions.

- The basic principle is that you may use Bindex in the same environment as your computer.
- Bindex is intended to be operated indoors, in clinics, hospitals or home healthcare use environments (see Appendix: Guidance and manufacturer's declaration - Electromagnetic Compatibility for power supply requirements) by a trained operator
- Bindex is not recommended to be used near active high frequency surgical equipment or in an RF shielded room of an ME system for magnetic resonance imaging, where the intensity of electromagnetic disturbances is high.
- Bindex is powered from the USB port of your computer. Please make sure that your computer is connected to a power source (battery or a mains outlet).
- Do not use Bindex near heat source or air conditioner and do not expose device to excessive moisture, above or under specified limits in section 5.
- Do not store your Bindex in a place where it exposes to sunlight.
- Measurements can be done while patient is either sitting or lying on a bed.

Computer hardware requirements					
Operating System:	Windows 8.1 or Windows 10				
Processor:	2 GHz, 32-bit or 64-bit				
Memory:	2 GB				
Hard Disk Drive:	Installation: 72 MB Bindex Standalone Software 353 MB LabVIEW Run-Time Engine 2012 109 MB Microsoft Access database engine 2010 2 MB device drivers				
	in use: 1.7 MB per patient NOTE: Bindex Standalone Software cannot be used if the remaining disk space is under 1 Gb.				
Screen Resolution:	1024x768				
Other:	USB port (type A) .NET framework v.4.0.30319 or newer (upgrade with Windows Update, if necessary)				

7. Setup

7.1. Unpacking Bindex

When you have received your Bindex BI-2 device package, remember to check that you have all components/parts which are listed in the packing list. Also remember to check that the packing list includes everything you have ordered. The package includes the Bindex BI-2 device and a Bindex BI-41 Measure. The electronic instructions for use (the User Manual) are stored in the accompanying USB drive. Additional product info card may also be included depending on market area.

The installation files for Bindex Software or a web installer application for downloading and installing Bindex Software are delivered on the accompanying USB drive or through a download link (provided by customer support).

7.2. Software installation

Installation of the Bindex Standalone Software should be done by a person with at least basic knowledge and experience about the Windows OS and installing new programs and hardware, e.g. a nurse with experience on the subject or an IT support technician. Software can also be installed by a Bindex representative as agreed.

PLEASE NOTE: Administrator rights are required for the installation.

To start the installation, start the web installer application and wait for the download to finish. Alternatively, if the install files have been delivered separately double click on the "setup.exe" file. You need to confirm that the program is allowed to make changes to the computer. You may also need to enter the administrator password before the installation launches. You can stop the installation at any time by pressing the CANCEL button in the lower right corner of the installation window.

Set the installation directories in the following window (Figure 4). Please verify

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the installation folder because you cannot move the software folder after installing.

The next step is accepting the license agreements (Figure 5). Carefully read the license terms before proceeding. Selecting "I accept the License Agreement" is required to use the Bindex Standalone Software. Press NEXT to continue. The next window includes the license terms for the National Instruments software required for Bindex Standalone Software. Accept the terms and click on NEXT.

Browse	1	
Browse]	
	Browse	

Figure 4: Setting the installation directories. The paths may be changed if needed.



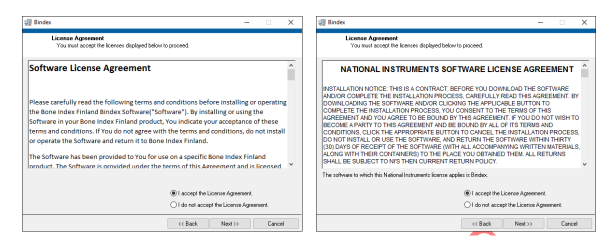


Figure 5: The license agreements. Carefully read the terms before proceeding. After accepting the licenses, continue with the installation by clicking on NEXT.

The following window (Figure 6) shows that you are about to install or change files related to Bindex Standalone Software. The installation starts by pressing NEXT.

After a successful installation a confirmation window is shown (Figure 7). Finish the setup by pressing NEXT. LabVIEW Run-time Engine 2012 and Microsoft Access database engine 2010 are required to be installed to run Bindex Standalone Software. The installation of these programs is automatically launched after the Bindex installation. Please follow the instructions on the screen to complete the setup. A restart is required before running Bindex Standalone Software. You may do this at this point or later.

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	allation. Ulick the Back but	ton to change the	Installation settings	i.		
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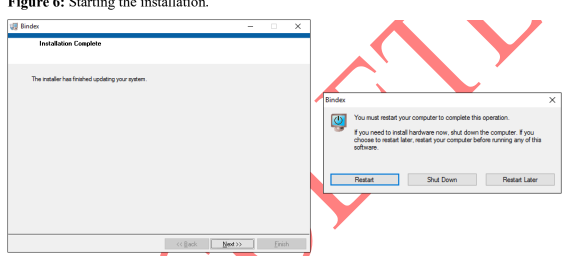


Figure 7: Installation completed successfully. Exit the installation program by pressing NEXT. After installing all required components, a restart is needed before using the software.



7.3. Device driver installation

Bindex BI-2 device drivers are installed together withBindex Software. After completing procedures in section 7.2, the Bindex BI-2 device may be plugged in for finishing the driver installation. Drivers are required for the computer to identify the Bindex device and to conduct the measurement correctly.

If the Bindex BI-2 device is plugged in before running the setup of Bindex Software and the computer is connected to the Internet, the operating system will automatically search for device drivers (shows up as FT240X USB FIFO) and install them (Figure 8). Please wait until a message about USB Serial converter having been successfully installed is displayed. If device drivers have been installed through Windows Update, it is recommended to also run the Bindex Software install. This is to make sure the latest compatible drivers approved by Bone Index Finland are installed.

After this Bindex Standalone Software is ready to be started.

X Device Setup	X Device Setup
Installing device Please wait while Setup installs necessary files on your system. This may take several minutes.	Installing FT240X USB FIFO Please wait while Setup installs necessary files on your system. This may take several minutes.
Close	Close

Figure 8: Device driver installation.

If the automatic installation is unsuccessful for some reason, a driver installation package is supplied with the installation files for manual installation. Please consult your Bindex representative or the Bindex Support and Service (see section 12 Contact information) for additional assistance.

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7.4. Software activation

7.4.1. First time activation

To start using the Bindex device you need to activate the software using an activation file. You will receive this file after sending a corresponding customer file to Bone Index Finland.

- Run Bindex Standalone Software (double-click on "Bindex.exe" in program folder or shortcut in your desktop). A dialog appears, asking you to save the customer file (Figure 9).
- 2) Save the file ("customer.key") to a folder of your choice, e.g. Desktop.
- 3) Send your customer file as an email attachment to info@boneindex.fi together with your contact information and the number of pre-paid analyses (PPA) to include. The expiration date of PPAs will be set according to the currently valid agreement between the customer and the seller.
- 4) An activation file ("activation.key") will be sent to you by email.



INFORMATION SECURITY HAZARD:

WARNING: Verify the email recipient carefully before sending files.

PLEASE NOTE: DO NOT edit the customer file or the activation file or they will become void.

Until the license has been activated, Bindex Standalone Software will ask for the activation file (Figure 10). After opening the correct activation file, software is ready for use (Figure 11).

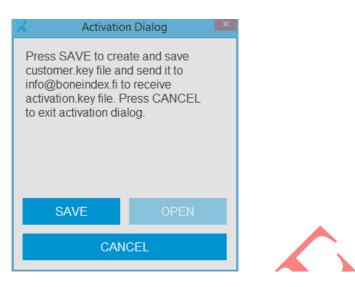


Figure 9: When you run the software for the first time a dialog appears.

Activation Dialog	
Press SAVE to create and save customer.key file and send it to info@boneindex.fi to receive activation.key file. To activate press OPEN and select activation.key file or press CANCEL to exit activation dialog.	
SAVE OPEN	
CANCEL	

Figure 10: Before the software is activated a dialog showing your customer key and asking for the activation key appears.

\mathbb{X}		×
	Activation succeeded!	
	ОК	

Figure 11: The software notification when the correct software activation file is opened.

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PLEASE NOTE: The database MUST NOT BE COPIED, or it will lock and require a new activation. In case of problems please contact your local distributor or Bone Index Finland Ltd. for assistance.

7.4.2. Ordering additional analyses

After activating the software each new Density Index analysis reduces the prepaid analysis counter by one. New measurements can be made until the PPA counter in the **Settings** page (Figure 28) reaches zero (all pre-paid measurements have been used). After this, software can be started and previous measurements viewed but no modifications can be made.

Additional PPAs can be ordered as follows:

- 1) Go to the Settings page and click on LICENSE AND PPA (Figure 12).
- 2) The **Order Dialog** opens. Click SAVE to create a new order file ("order.key") and save it to a location of your choice (e.g. Desktop).
- 3) Send the order file as an email attachment to info@boneindex.fi with your contact details and number of PPAs to order.
- 4) Additional PPAs will be sent to you in a PPA key file ("ppa.key") in return.

INFORMATION SECURITY HAZARD:

WARNING: Verify the email recipient carefully before sending files.

To activate the additional analyses:

- 1) Select OPEN in the Order Dialog and locate the "PPA.key" file.
- 2) Select and open the PPA file.
- 3) A notification about successfully adding PPAs and their number will be shown. After clicking OK, the new PPAs are immediately usable.

Each activated PPA package can be viewed by clicking on SHOW PPAs in the **Order Dialog**. The PPA packages are depleted and listed in the order of their expiration date, starting from the first package to expire. Depleted and expired packages are removed from the list automatically.

Operator: First Lastname Operator select: Last operator Language: English Units: Metric Pdf directory: C:\Program Files (x86)\B Reference DXA manufactu GE	CANOFI	Company logo: X I I N D E X I Sent:	Vorder Dialog Pres orde info(PPA PPA PPA pres orde orde CLOS	X date 31.12.2017 y SE WINDOW
---	--------	--	---	---

Figure 12: License and PPA window. You can create new order keys and add ordered PPAs from this menu. A list of activated PPA packages can be seen by clicking on SHOW PPAs.

7.4.3. Additional software

You can export the result sheets as PDF files. To be able to view them a PDF reader application (e.g. Adobe Reader) is required.

The User Management Application (UMA) is required for creating and managing Bindex users and setting the password protection for Bindex Standalone Software. UMA is automatically installed with the Bindex Standalone Software standard installation. For more information, see section 7.5.

7.4.4. Additional system setup

Adjust the computer screen brightness to a comfortable level from the system/computer settings. Too low screen brightness may hinder the use experience and the measurement in Bindex Software.

Enable the system sounds of the computer to hear the audio signals for accepted signals during Bindex measurement.

For high resolution displays (above Full HD, e.g. 4K) it is recommended to use the Windows scaling function (Settings \rightarrow Display \rightarrow Scale and layout). Set the scaling to a level at which the Bindex Software window can be viewed with ease.

7.4.5. Information and system security

For ensuring data security it is highly recommended to use password-protected Windows user accounts or other authorization method for the computer Bindex Software is installed on. By default the password protection and user accounts for Bindex Software are on. Please see 7.5 User Management Application (UMA) for additional information.

Up-to-date antivirus, malware protection and a firewall (Windows Defender firewall) is also highly recommended for protecting the patient and operator data. It is recommended to enable automatic updates for the Windows operating system, firewall and virus/malware protection.

Bindex Software does not need connection to internet. However, most computers are usually connected to the internet when in normal use. Therefore it is vital to ensure the safety of the network environment when using Bindex Software. As a general rule connect only to secure local networks. Avoid using unsecure public networks.

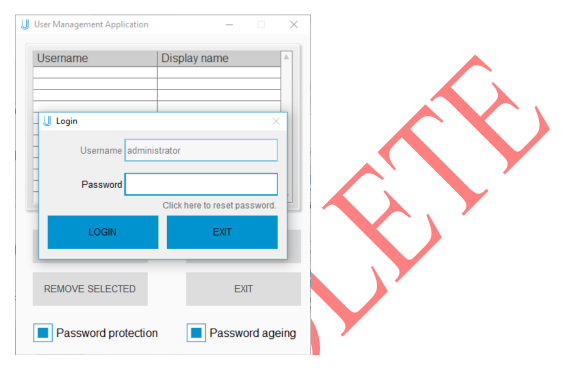
In Bindex Software all patient and measurement data is stored in an encrypted format. Data exported from the program (database exports and PDF reports) is not encrypted by Bindex Software. The customer must ensure the exported data is stored in a secure manner to prevent unauthorized access.

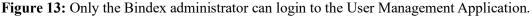
Bindex Software does not use automated backups. Please make sure the Bindex database and measurement files (by default located in the folder C:\Program Files (x86)\Bindex\database) as well as all other important files are backed up using e.g. Windows File History to a secure location regularly. Bone Index Finland is not responsible for any loss of data on a measurement computer.

Always follow your organization's IT guidelines and security practices. In case of questions, please contact Bindex Support and Service (see 12 Contact information).

7.5. User Management Application (UMA)

Bindex Standalone Software users are managed in a separate program, the User Management Application (UMA). UMA is installed in the Bindex main folder and is started automatically upon first startup of Bindex Standalone Software. It can later be used by double-clicking on "UMA.exe" file.





Only the "administrator" account can login to UMA. The administrator can create a needed number of operator accounts for Bindex Standalone Software. At least one operator account needs to be created to access Bindex Standalone Software, as the administrator account cannot be used in measurement software. The first login is made using the password "changeme". After entering the password, the administrator is required to change the password before continuing (Figure 14). The password is required to be at least eight (8) characters in length and contain at least one uppercase and lowercase letter, a number and a special character from the following:

~! @ #\$%^&*_-+='``|\() { } []:;.,"<>?/

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U New password		\times
Enter new pass	word for administrator.	
New password		
Confirm password		
СН	ANGE PASSWORD	

Figure 14: The administrator is required to change the password after the first login.

When logged in, the currently active users are shown in a list displaying their:

- 1) Username (used only for logging in to Bindex Standalone Software)
- 2) Display name (shown in PDF reports, e.g. full name or anonymous JD)

	listrator tor Two	
INSERT OPERATOR	MODIFY SELECTED	
REMOVE SELECTED	EXIT	

Figure 15: User Management Application main window.

When ready, click on EXIT to close UMA.

7.5.1. Password protection

Using operator accounts and password protection is on by default. The feature can be enabled or disabled by using the PASSWORD PROTECTION checkbox.

When the option is disabled, any person with access to Windows and Bindex Standalone Software can conduct and view patient measurements. The locking functionality is also disabled in Bindex Standalone Software.

The PASSWORD AGEING option allows the administrator to enforce the users to regularly change their password to Bindex Software. When enabled, six months after the initialization of the last password the operator is required to change their password upon login to Bindex Software.

7.5.2. Creating new operators

New operators are added to Bindex Standalone Software by clicking on INSERT OPERATOR. Enter the *Username*, initial *Password* and *Display name* for the new operator (Figure 16).

📙 Insert operator		Х
Username:	operator	
		_
Password:	3P}?sl5f	
Display name:	Operator One	
Diopia y name.		
PANDOM	ZE PASSWORD	
KANDOW	ZE FASSWORD	
CREATE USE	R CANCEL	

Figure 16: Creating a new operator account.

The *Password* is required to be at least eight (8) characters in length and contain at least one uppercase and lowercase letter, a number and a special character from the following:

~! @ #\$%^&*_-+='``|\() { } []:;.,"<>?/

The password can also be randomized by clicking on RANDOMIZE PASSWORD, suggesting a password with 8 random characters. The initial password must be changed by the operator after first login.

When finished, click on CREATE USER to save the new operator to the Bindex database. Press CANCEL to stop the operator creation at any time.

7.5.3. Modifying operator details

Operator details (Password and Display name) can be edited by selecting an operator from the list and clicking on MODIFY SELECTED. The window showed in Figure 17 opens.

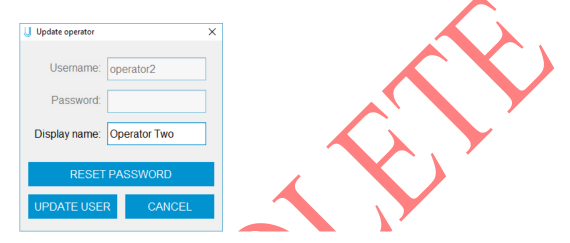


Figure 17: Modifying operator details. The administrator can change the Display name or e.g. reset a forgotten password of an operator.

The *Display name* of the operator can be changed by setting the name and clicking on UPDATE USER.

To modify the password (e.g. if forgotten by the operator), click on RESET PASSWORD and enter the new password. See section 7.5.2 for password strength requirements. The RANDOMIZE PASSWORD button becomes available after resetting the password. Click on UPDATE USER to save the changes.

Click on CANCEL at any time to revert the changes and return to the main window.

7.5.4. Deleting operators

Operator accounts that are no longer needed can be removed from the database. After deleting them through UMA, the account holder can no longer login in Bindex Standalone Software. The operator data is still saved in previously made measurements.

To delete an operator account, select the operator from the list in the main window and click on REMOVE SELECTED. A confirmation window is displayed (Figure 18). Answer YES to confirm the deletion of the operator account. Select CANCEL to cancel the operation.

User Management Application	- 🗆 X	
Username administrator operator2	Display name administrator Operator Two	
Do you want to de	elete user 'operator2'?	
INSERT OPERATOR	MODIFY SELECTED	
REMOVE SELECTED	EXIT	
Password protection	Password ageing	

Figure 18: Confirmation for deleting an operator account.

PLEASE NOTE: After deleting an account, the username is still permanently reserved. Another operator account with the same username cannot be created.

PLEASE NOTE: The "administrator" account cannot be deleted.

7.5.5. Resetting the administrator password

If the administrator forgets the password, it can be changed by clicking on

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CLICK HERE TO RESET PASSWORD text in the login window (see Figure 13). A dialog window opens, displaying a *Request Key* (Figure 19). To reset the password, follow these steps:

- 1) Copy the *Request Key* to an email message.
- 2) Send the key along with a request to change the administrator password.
- 3) The Reset Key will be sent by email.
- 4) Copy the Reset Key to the corresponding field in UMA and click RESET.
- 5) The program informs about successfully resetting the password.
- 6) Enter the new administrator password twice and click on CHANGE PASSWORD to continue (see Figure 14).

U Reset password	×	
Send the <i>Request key</i> to Bone Index Finland to receive the <i>Reset key</i> .		
Request Key: 73737373		
Reset key:		
RESET CANCEL		

Figure 19: The dialog for resetting the administrator password.

In case of any issues with accounts or passwords, please contact Bindex Support and Service (see 12 Contact information).

7.6. Bindex device setup

After completing the software and driver installation (see 7.2 and 7.3), the device is ready to be used. Connect the Bindex device to a free USB port (type A) of your computer. To test that the device has been correctly installed:

- 1) Start Bindex Standalone Software and log in.
- 2) Click on OPEN CASE and search for "test patient".
- 3) Select the test patient and click OPEN.
- 4) Conduct a calibration as instructed in 8.6 Bindex quality verification. If the calibration is successful, the device has been correctly installed.

Once connected to a USB port, the device is in stand-by state.

PLEASE NOTE: The ultrasound transmission is on only during a measurement in Bindex Standalone Software.

It is recommended to disconnect the Bindex when not in use to reduce the power consumption of the computer. If kept connected continuously, the Bindex device may feel slightly warm to the hand, but this is harmless. The device is not equipped with an external power switch or button.



GENERAL SAFETY HAZARD:

Do not modify this equipment without a specific authorization by the manufacturer. A modified device cannot be guaranteed to comply with the requirements of the applicable safety standards and regulations.



ELECTRICAL HAZARD:

Do not use a multiple socket outlet to connect the system to the power grid. The electrical safety of the system cannot be guaranteed for all required fault conditions if a multiple socket outlet is used.



ELECTRICAL HAZARD:

Use a IEC 62368-1:2014 compliant isolation transformer to connect the measurement computer to the power grid. (Not needed when the computer is used with a battery). The electrical safety of the system cannot be guaranteed in case a non-compliant power source is used.

8. Using Bindex

8.1. Connecting and disconnecting the Bindex device

Connect the Bindex device into the USB port of the computer before starting the pre-measurement calibration. This can be done before or after launching Bindex Standalone Software. If the device is disconnected from the computer while the software is running, the connection can be restored by reconnecting the device and pressing CALIBRATE in the **Measurement** window.

Operation can be safely terminated by shutting down the software by pressing the "X" at the upper right corner of the window and unplugging the pulser unit. If a patient case is open, a confirmation will be prompted before exiting. In case of an emergency, operation may be stopped by just unplugging the USB cord while the software is running, but data loss may occur.



ELECTRICAL HAZARD:

Do not touch the connectors of the computer and the patient at the same time. Touching the connectors of the system results in potential electrical safety hazard. Where possible, the computer should be kept away from the patient area.

8.2. Basics of Bindex Standalone Software

PLEASE NOTE: Bindex Software cannot be used with a Windows Guest account. Please use a Local user or Administrator account.

Bindex Standalone Software is started by double-clicking on "Bindex.exe", the Bindex shortcut created on the Desktop or by selecting "Bindex" in the Windows Start Menu.

Depending on the password protection setting in User Management Application (UMA), the login will be made as described in 8.2.1 or 8.2.2.

8.2.1. Logging in with password protection enabled

If password protection is enabled in UMA, at least one user must be created before logging in to Bindex Software is possible. The "administrator" user cannot be used for logging in.

💥 Login	×
	_
Enter username and password.	
	- 1
Username	
	- 1
Password	
LOGIN EXIT	

Figure 20: The Login window of Bindex Standalone Software, when password protection is enabled.

At startup, a login screen will be displayed, asking for the *Username* and *Password* (Figure 20). Enter the username and password to their respective fields and click on LOGIN. The operator is asked to change their password at first login (Figure 21). Enter the previous password once and the new password two times and press CHANGE to move on to the front page of Bindex Standalone

Software.

💥 Login	×
Enter new passwo	ord.
Username	operator
New password	
Confirm password	
CH	IANGE PASSWORD

Figure 21: At first login, the user is required to change the password.

8.2.2. Logging in with password protection disabled

If password protection has been disabled in UMA, at startup the software asks for the operator name (Figure 22) or alternately uses a name defined in your settings (see 8.2.3.3 Settings). Enter the operator name to use and click on LOGIN to move on to the front page of Bindex Standalone Software.

Enter operator name	
Enter operator name	
EXIT	LOGIN

Figure 22: The Login view, when password protection is disabled. Press the LOGIN button to set the operator name and continue to the front page of the program.

8.2.3. Front page and functions

There are four buttons on the front page (Figure 23): NEW CASE, OPEN CASE, SETTINGS and ABOUT.



Figure 23: Front page of Bindex[®] software. You can always get back to this page by pressing the HOME button in the upper right corner.

When a patient case is open, the tabs **Patient**, **Location**, **Measurement**, and **Results** on the upper bar can be used to quickly navigate between measurementrelated data. When no case is open, the **Patient** tab can be clicked for creating a new patient case.

Click on the HOME (symbol) button to return to the front page at any time.

The "?" (symbol) button opens the User Manual in a PDF reader in the language chosen in **Settings**. If there is no translation for the selected language, the English version is shown.

The LOCK (symbol) button is visible when password protection has been enabled in UMA. Pressing the button at any time locks Bindex Standalone Software and prevents its use until the current user logs in again or EXIT is pressed (Figure 24).

Bindex Standalone Software locks the program automatically after being idle for 30 minutes. The automatic locking is opened similarly to the manual locking (user logs in again or exits the program).



PLEASE NOTE: Exiting Bindex Standalone Software when the program is locked causes all unsaved data to be deleted!

	💥 Unlock software	×	
	Software is locked. Unlock or exit the software. If software is closed all unsaved data will be lost		
	Username operator	5	
	Password		
	UNLOCK EXIT		
/	H. C.		

Figure 24: After 30 minutes of inactivity Bindex Standalone Software is locked. The program can also be locked at any time by pressing the LOCK (symbol) button.

8.2.3.1. New case

Press NEW CASE to start a measurement with a new patient (see 8.3 Patient information). If you have a previous case open you will be asked a confirmation to proceed (Figure 25). Selecting OK starts a new empty patient case but all unsaved data from the current measurement is lost.

If the patient's data has been saved previously, please use the OPEN CASE for a new measurement.

Ж	×
	is already open. o start a new case?
Yes	No

Figure 25: The software asks a confirmation before starting a new case over an existing case.

8.2.3.2. Open case

Use the OPEN CASE page (Figure 26) to:

- 1) Add another measurement to a previously measured patient.
- 2) View the results of a measured patient or make corrections to their details.
- 3) Delete patients or measurements.
- 4) View all the measurements of a single patient in timeline view.

A patient can be searched by typing the first or last name or patient ID on the *Search term*. Separate the used search terms (up to three) with spaces. Bindex Software automatically suggests patients starting with the letter or phrase typed in this field.

The patient list columns (last name, first name, PID) can be sorted in ascending order by double-clicking on the title. After selecting a column for sorting, the title appears as bolded.

Select a patient and press OPEN to start a new measurement and continue to the **Patient** page which shows the selected patient's previously saved details. You can also double-click on the patient list to start a new measurement for the selected patient.

To view previous results, select a name and measurement from the lists and press the RESULTS button. The **Results** sheet of the previous measurement opens. You can also double-click on the result list to view the results for the selected measurement.

Select TIMELINE to see all results of the selected patient in a timeline view if the patient has been measured more than once. The timeline can also be exported as a PDF file.

The DELETE button can be used to delete either the selected patient or the selected measurement (Figure 27). Select MEASUREMENT to delete only the selected measurement from the selected patient. Select PATIENT to delete the entire patient file including all the measurements.

The patient and measurement data can be exported to a text file by clicking on EXPORT DB. The exported content can be limited by filtering the patients by date of birth or measurement date. Press EXPORT ALL to export all measurement information currently saved in the database.

PATIENT		MEASUREMENT	DEO	11 T C		0
PATIENT						
Search term: sam						
last name Patient	first i Sampl		PID 123-456		date	
				- III E		
					v	
	_					
DELE	E	EXPORT DB		RESU	JLTS	
BACI	<	OPEN		TIME	INF	
				8	🖁 вім	N D E

Figure 26: Open case view. By using the BACK button you will continue to front page of the software.

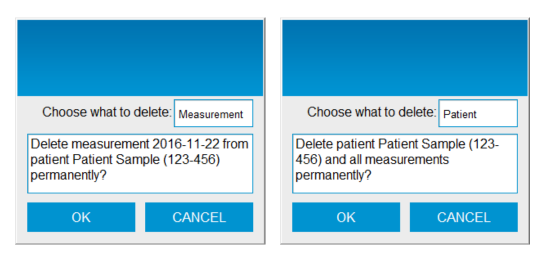


Figure 27: Delete a patient or measurement from the database.

8.2.3.3. Settings

By pressing the SETTINGS button, the operator naming method, software language and company information (contact details and logo) can be changed (Figure 28).

Click on the white box under COMPANY LOGO to select the image file to use. The supported image formats are JPG, BMP and PNG. The image is automatically fitted to the measurement report. Best quality is achieved with a square-shaped image (side ratio 1:1).

The number of available PPAs is shown at the bottom of the page in the info bar. The measurement counter since the day of the software activation is also shown.

For more information about PPA ordering, please see 7.4 Software activation.

Operator: Operator One Operator select:	Bone Index Finland Ltd.	
Operator select:		×
	Address - line 1:	BINDEX
Last operator	Savilahdentie 14	
Language:	Address - line 2:	
English	70700, Kuopio, FINLAND	
Units:	Phone number:	_
Metric	+358 50 448 1696	
Pdf directory: C:\Program Files (x86)\Bindex Reference DXA manufacturer: GE	Reports FRAX questions: Ask patient consent	Print signals:

Figure 28: Change settings view. In this page you can change the program, language, operator and company information settings and the default directory for saving the PDF reports.

When password protection is on, *Operator* and OPERATOR SELECT options are disabled. In this case the operator can be changed by closing the program and logging in with another user's details.

If password protection is disabled in the UMA, the operator name and the default operator name preset is selected (OPERATOR SELECT) can be changed.

- 1) LAST OPERATOR: the operator name is asked at startup, but the name of last operator is suggested.
- 2) WINDOWS LOGON NAME: software opens without asking the name of the operator. The Windows logon name is used.
- 3) LOCKED: software opens without asking the name of the operator. The last saved operator name is used.

The language of the software interface can be chosen in the **Settings** page. The measurement units (Metric, Imperial or U.S.) to be used with patient information can also be chosen.

The default location for saving the PDF reports can be set by using the PDF DIRECTORY control. The saving location can still be set separately for each report.

You can select which manufacturer's DXA device values to use as reference for calculating the BMD approximation by selecting a manufacturer from the dropdown menu REFERENCE DXA MANUFACTURER. This will slightly modify the BMD thresholds in accordance with the manufacturer in question.

When FRAX QUESTIONS option has been enabled, a FRAX questionnaire will be shown after the patient info has been entered (see 8.3 Patient information) When the selection is disabled, only the basic information of the patient is entered before measuring.

The ASK PATIENT CONSENT option adds a checkbox to the **Patient** tab, when enabled. The checkbox needs to be ticked before saving any patient information to the database. The checkbox acts as a reminder if local legislation or regulations require certain information to be provided to or permissions acquired from the patient before saving their personal information to the system.

The PRINT SIGNALS option allow the operator to included the measurement signals as images to the measurement PDF report. When enabled, the signals from the five repetitions are added to the report. The signal peak locations can also be seen in the images. When disabled, the signals and the peak locations can still be viewed in the Results tab of Bindex Software using the VIEW SIGNALS button.

CHANGE PASSWORD button is visible, if password protection is enabled in UMA. Click on the button to open a dialog (see Figure 29). Enter the previous password once and the new password two times. Click on CHANGE to change the password. Please see section 7.5.2 for password strength requirements. A confirmation message is shown when successful.

X Change password ×				
Enter old password and new password twice to change the password.				
Old password				
New password				
Confirm password				
CHANGE	EXIT			

Figure 29: Changing the operator password. Enter the previous password once and the new password twice, then click on CHANGE.

The LICENSE AND PPA button is used when additional analyses need to be ordered. Please see section 7.4.2 Ordering additional analyses for more information.

8.2.3.4. About

The ABOUT button will open the Bone Index Finland Ltd. web site (https://www.bindex.fi) in your default browser.

8.3. Patient information

On the **Patient** page (Figure 30), the name, ID, date of birth (DOB), sex, ethnicity, weight and height of the patient are entered. In addition, a comment field is located at the bottom of the page for entering any information that should be included in the results printout (e.g. risk factors for osteoporosis). The SAVE button saves the patient info to the database for later use. The NEXT button will lead to the next page (**Location**). Before saving or continuing, the "Consent obtained" checkbox needs to be ticked, if it has been enabled in the **Settings** (see 8.2.3.3 Settings).

If an existing case has been opened and patient information is changed, you will be asked (Figure 31) whether you want to update the data or discard the changes.

If you save or try to continue to the next page before you have filled patient information you will be notified to do this (Figure 32).

PATIENT	LOCATION MEASUREMENT	RESULTS		
1. First name	Sample		SAVE	
2. Last name	Patient			
3. ID 4. Date of birth	123-456 1.2.1953 ID Age 66			
5. Sex	Female			
6. Ethnicity	Caucasian		Consent obtained:	
7. Weight 8. Height	75 kg 170 om			
9. Comments				
To save or update pa	tient info press SAVE, to continue press NEX	ſΤ	💥 BINDEX	

Figure 30: The Patient page. All information must be entered before you can continue by using the NEXT button. By saving the patient info you can find the info from OPEN CASE later.

PLEASE NOTE: The weight and height of the patient should be measured if possible! This way you will get the most reliable measurement result.

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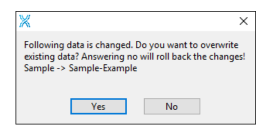


Figure 31: Software asks confirmation before writing over existing data.

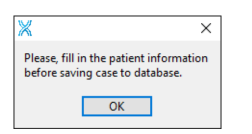




Figure 32: You will be asked to fill the necessary patient information before continuing to the measurement.

If enabled in Settings, the FRAX questionnaire will be shown after clicking NEXT in the Patient page (Figure 33). This questionnaire can be used to save the information of the (online) FRAX test. The information is also printed in the PDF report.

Bindex Software DOES NOT conduct the actual FRAX analysis or give out the recommendation but can only be used to save the input info and results.

The data entered in the FRAX questionnaire DOES NOT affect the Density Index calculation.

PATIENT LOCATION		000
1. Previous fracture	No Yes	
2. Parent fractured hip		
3. Current smoking		
4. Glucocorticoids		
5. Rheumatic arthritis		
6. Secondary osteoporosis		
7. Alcohol 3 or more units per day		
8. BMI, kg/m²	0,0	
9. Major osteoporotic	0,0	
10. Hip fracture	0,0	
11. FRAX recommendation	Undefined	
Fill in the questions and DXA results and pr	ess NEXT to continue	💥 BINDEX

Figure 33: If enabled in settings, the FRAX questionnaire can be filled in after the basic information of the patient has been entered.



8.4. Patient positioning

For the duration of the measurement, the patient should be lying on e.g. a bed. Alternatively, the patient may be sitting and the examined leg is straightened and supported with e.g. a chair. Find a comfortable position for yourself and the patient (Figure 34). Clothing must be removed below the knee up to over the ankle.



Figure 34: Patient positioning on a bed. Remember to keep an ergonomic position when you are measuring.



8.5. Measurement site location

By default, the measurement is conducted on the left tibia. When selecting the measurement location, remember that the Bindex measurement is contraindicated for locations with:

- broken or irritated skin;
- a fractured bone;
- implants, plates or fixations.

In case of contraindications on only the left tibia, conduct the measurement on the right tibia.

To determine the right measurement location, use the Bindex Measure. Before using the Measure you need to locate the upper head of tibial bone (the knee joint) (Figure 35). It may be helpful to move the patient's leg while palpating the knee joint. When you have located the knee joint, mark it on the skin.

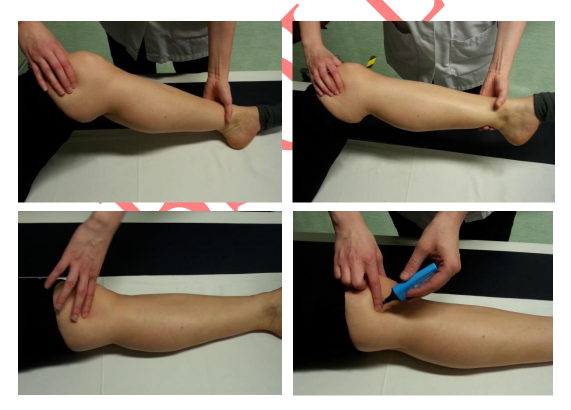


Figure 35: Locating and marking the knee joint. First you must locate the upper head of the tibia or the knee joint.

Put the arrow head of Bindex Measure on the distal head of tibia (on the medial

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malleolus, see Figure 36). Check the number on the scale 1 (or A) at the mark on the knee joint (e.g. number 12 in the Figure 37). After this, find the same number on the scale 3 (or C) (mark this number as in Figure 37). This is the measurement site.

This site is 1/3 of the length of the tibia from the upper head (Figure 37 and Figure 38). Please enter this number to in the **Location** page of Bindex Standalone Software (Figure 39). After this click on the NEXT button to continue.



Figure 36: Locating the distal head of tibia. The arrow head of the stick is located on the medial malleolus. After this, check the number on the Bindex Measure at the mark at the knee joint.



Figure 37: Locating the measurement location. The right measurement location can be found at the same number on scale 3 (or C), e.g. number 12 in this picture.



Figure 38: The tibia typically has a plate-like cortical surface at this site. The measurement should be made at the center of cortical bone plate.

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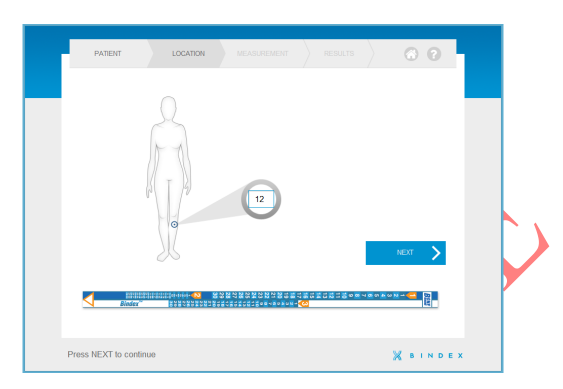


Figure 39: Enter the measurement location number from the Bindex Measure in the Location tab.



8.6. Bindex quality verification

Every time, the first thing to do on the measurement page is the quality verification check with Bindex. In the calibration the echo from the surface of the transducer is analyzed in order to verify that your Bindex is working properly.

Let the transducer head be freely in air (Figure 40) and press the CALIBRATE button (Figure 41). Make sure that the head of transducer is clean and there is no gel on it. The Bindex software shows a message if the calibration was not successful (Figure 42).

If the calibration fails, please check that the probe surface is clean and that the probe is properly connected to the computer before trying again. If the device has been kept in a hot or cold environment, the device may need to settle closer to room temperature (operating temperature 15-40°C) before the calibration succeeds. If the calibration fails consistently, please contact your local distributor or Bindex Support and Service for assistance.



Figure 40: The transducer head should be freely in air when you press the CALIBRATE button.

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	S	BOOST	
		DISCARD	
	G	Messuring	
Noise level	~~		

Figure 41: The CALIBRATE button. The button for calibration is located at the upper right corner. Measurement cannot be started before a successful calibration.



Figure 42: A failed calibration. Software notifies the user if the calibration was not successful.

8.7. Measurement with Bindex

To begin the measurement, first apply ultrasound gel on the skin over the measurement location. Turn on the measurement by pressing the circle button on the center of measurement page (Figure 43). The green segment on the circle lights and circular measure light turns to green under the buttons. Place the transducer on the skin beside the measurement location and move it slowly over your mark on the skin (Figure 43). When you clearly see two echo spikes in the signal window, you are at the right site. You may need to adjust the angle of the transducer to maximize the reflections.

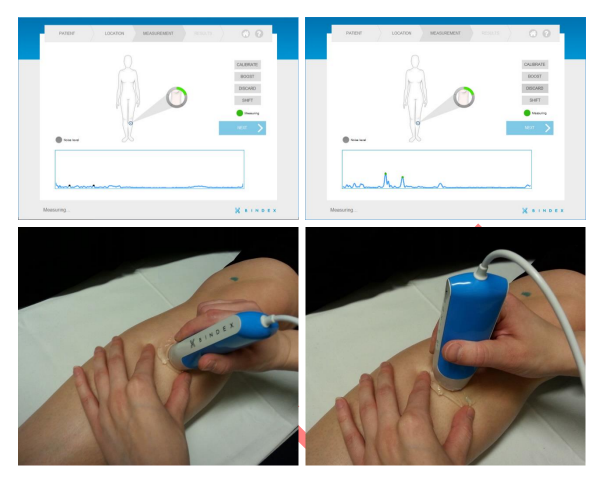


Figure 43: Conducting the measurement. First put the transducer next to bone and then move it over the bone. Keep an eye on the ultrasound signal on the signal window. When you see two echo spikes in the signal you are at the right location.

The software accepts the measurement automatically when the echoes are strong enough. The echo is strong enough when the dot marker on the echo peak turns from black to green. **Please note that observing the shape of the signal is also very important for a successful measurement!** For more information about signal acceptance please see 8.8 Signal Acceptance Window.

When a measurement is accepted an audio signal is played and the measurement stops for a short time (the signal is green at this time, the last accepted signal is shown). The transducer should be lifted off from the skin and the operator can proceed to make the next measurement similarly as explained above. If the signal is too noisy the indicator "Noise level" is lit on the left side of the screen. Adjust the transducer location or angle to reduce the noise.

The circular measurement button has five segments which will light green upon a successful measurement. When five measurements are accepted the green circle is complete, the measurement stops and the user is directed to the **Signal Acceptance Window**. Information on accepting and discarding measured signals can be found in 8.8 Signal Acceptance Window

PLEASE NOTE: If the echo spikes are too weak for acceptance you can add gain to the signal by using the BOOST function.

The measured signal can be amplified by pressing the on-screen BOOST button or Spacebar on the keyboard of your computer (Figure 44). Additional boost is removed when the measurement is restarted. Press the measurement circle again to start measuring again without boost. The Boost functionality should only be used if the signal peaks can be located reliably, but the amplitude is not high enough for acceptance.



ELECTRICAL HAZARD:

Do not touch the connectors of the computer and the patient at the same time. The patient must be separated from the measurement computer connectors to preserve the electrical safety of the system.



INCORRECT USE HAZARD:

Do not use the BOOST function before detecting and attempting to capture the echo spikes! Adding amplification before setting the transducer to the correct location may cause incorrect reflections to be accepted upon conducting the measurement.



INCORRECT USE HAZARD:

Overusing the BOOST function may cause incorrect reflections or noise to be interpreted as the echoes from the bone surface. Only add amplification if the measurement location can be confirmed to be correct.



Figure 44: The use of the BOOST button. Measurement signal before (left) and after (right) boost effect.

If the software accepts a signal which is too noisy (Figure 45) e.g. due to excessive amplification, you can delete the measurement by pressing the DISCARD button. The last accepted measurement is removed. Signals can also be discarded from the Signal Acceptance Window, as described in the next section. You can see the accepted signals later in the Results page by pressing VIEW SIGNALS (Figure 50).



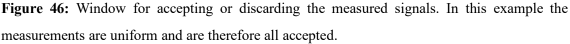
Figure 45: A noisy signal. If too much amplification is used, you may see a very noisy signal with multiple strong echo spikes. An acceptable measurement only includes two strong spikes.

Sometimes plenty of soft tissue or swelling at the measured limb may cause the echo peaks to be cropped out of the default signal window. By clicking on the onscreen SHIFT button the signal window is set to show a later time frame, i.e. echoes deeper from the tissue can be viewed. To the operator this appears as the signal plot being moved towards the left side of the window. The SHIFT button turns blue when activated.

8.8. Signal Acceptance Window

After the five measurements on a site the accepted signals from each repetition are shown in separate signal boxes (Figure 46). This enables you to monitor the quality of measurements and to remove faulty or inaccurate measurements from the series.





Signals that deviate significantly from the average of all measurements are marked with a red indicator at the top left corner of each signal box (Figure 47). A green indicator is shown at the upper left corner of the signal window if the signal is close to the average. The signals marked with a red indicator at the upper left corner are removed from the measurement series. If any signals are removed, software will return to the Measurement window.

The signal peaks used for thickness measurement are marked with green indicators.



You may select any number of signals to be discarded or accepted, even the deviating signals automatically marked by the software. The status of the signal box can be changed between discarded/accepted by clicking on the signal window. The color of the indicator changes along with the status of the signal. Examples of discarded signals and their possible causes are given in Figure 48 and Figure 49.

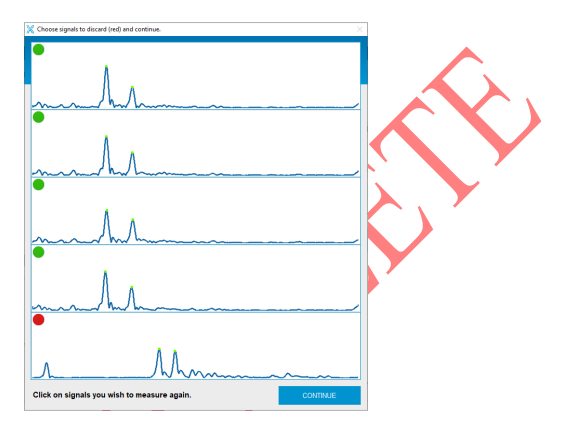


Figure 47: A signal deviating from the average. In this figure the bottom signal deviates significantly from the average of all measurements and is therefore suggested to be discarded by the software.



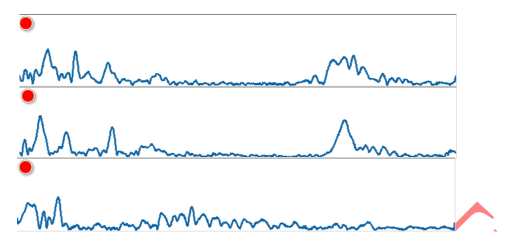


Figure 48: Excessive amplification. All signals shown in this figure should be discarded because they show too much noise and multiple high peaks due to the excessive use of the BOOST button.

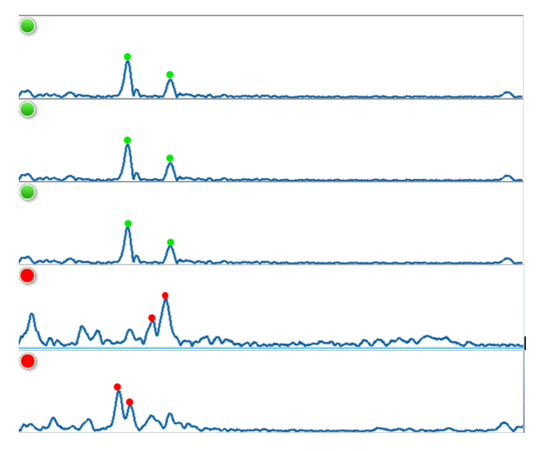


Figure 49: Incorrectly positioned probe. The lower signals marked as red deviate significantly from the average because the probe has been in a tilted position in comparison to the measurement location. The amount of noise generated in the signals is also high. The signal peaks accepted by the software have been marked to the figure with dots.

If one or more measurements are selected to be discarded, new measurements need to be made to replace the removed ones. In this case, after pressing CONTINUE the user will be guided back to the **Measurement** page. New measurements can be made after pressing the measurement circle. New calibration of the Bindex device is not required after returning from the signal selection window. When all five repetitions are accepted, the user is guided directly to the **Results** page.

It is important to check the uniformity of the measured signals always before accepting them. The most important factors to monitor in the signal acceptance window are:

- 1) the distance between the signal peaks
- 2) the location of the signal peaks
- 3) the shape of the signal.

If any of these factors deviates significantly from the others in one or two measurements, the deviating signals should be removed from the measurement series and new measurements should be conducted. At least three signals should be similar to each other. Otherwise, the series should be re-measured (turn all status indicators red and press CONTINUE).

8.9. Interpretation of the Bindex results

On the results page you will find the Density Index value which is an estimation of total hip bone mineral density measured with DXA (Figure 50). The value is also presented as a marker (blue arrow) on a three-color scale (green, yellow and red). The marker on the green zone indicates a "Low Probability of Osteoporosis". The marker on the red zone indicates a "High Probability of Osteoporosis". If the marker is in the yellow zone, the patient needs additional investigation for the determination of osteoporosis status.

If the marker is off the visible scale (under 0.6 or over 1.2 g/cm²), a notification is displayed. The user is prompted to check the patient details for possible typographical errors, which may cause atypical results.

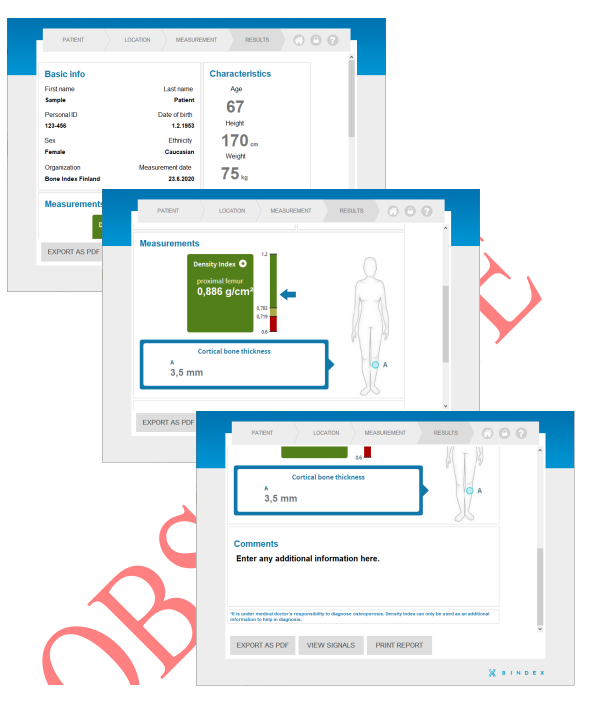


Figure 50: The result page includes patient information, Density Index and apparent cortical bone thickness values. In addition, the Density Index value is also presented on a three-color scale (green, yellow and red).

The results can be exported to PDF format by clicking EXPORT AS PDF and choosing the location for saving the file. The report can also be printed by clicking PRINT REPORT and given to the patient. Please go through the information shown in the report together with the patient first.

PLEASE NOTE: It is under the Medical Doctor's responsibility to diagnose osteoporosis. Density Index can only be used as additional information to help in diagnosis.

The color scale is based on a 90% sensitivity and specificity threshold analysis which has been determined in the clinical trials. The threshold value 0.84 between the green and yellow zone presents the sensitivity threshold and value 0.78 between the yellow and red zone present the specificity threshold. For Bindex the thresholds for osteoporosis with 90% sensitivity and specificity have been determined in population with 75 osteoporotic and 373 healthy patients (Karjalainen et al. Osteoporos Int 2016).

Originally the use of thresholds was proposed by Blake et al. Osteoporos Int Issue:16; 2005. It was suggested that the thresholds should be determined with at least 70 osteoporotic and 70 healthy subjects. Thereby, it is ensured with 95% confidence that the true sensitivity and specificity of a method will not fall below 80%.

The Density Index is a predictive index of proximal femur bone mineral density and thus intended for detection of hip osteoporosis. The apparent cortical bone thickness reported by Bindex has been assessed from measurement of the time difference between the echoes from periosteum and endosteum and using estimated constant speed of sound. The speed of sound in bone may vary due to the biological changes in bone tissue and structure. Therefore, the reported cortical bone thickness is an estimate. The apparent cortical bone thickness is an important input parameter into the Bindex for the determination of Density Index.

9. Cleaning, disinfecting and packing

The parts of the Bindex system that come into skin contact with the patient (i.e. the surface of the transducer, the handpiece and the BI-41 Measure) should be **cleaned (or checked for cleanliness) and disinfected before each patient measurement**. This is to minimize the risk of cross-infection between patients.

It is recommended to use isopropanol disinfective solutions (up to 70%) designed to be used on ultrasound transducers (for instance Transeptic, Parker Laboratories Ltd.). Ethanol solutions must not be used. Protective gloves are recommended during the cleaning and disinfection to minimize skin irritation.

Precautions for preventing damage to the Bindex device upon cleaning and disinfection:



MECHANICAL HAZARD:

DO NOT allow sharp or hot objects to touch the transducer or the cable.



MECHANICAL HAZARD:

DO NOT bump the device on hard surfaces while handling.



ELECTRICAL HAZARD:

DO NOT immerse the device in water or cleaning liquids to prevent the accumulation of liquid inside the device. Bindex is not intended to be immersed. The device can be sprayed lightly to moisten its surface.



MECHANICAL AND ELECTRICAL HAZARD:

DO NOT USE cleaning solutions that can damage the plastic of the device, such as ammonia, acetone or strong acids. If unsure, please check the suitability from the manufacturer of the solution before using it.



MECHANICAL HAZARD:

The device MUST NOT BE disinfected or sterilized using heat or steam, e.g. in an autoclave. The device does not need to be sterilized; regular disinfective solutions are sufficient to guarantee patient safety.



MECHANICAL HAZARD:

DO NOT USE regular rough tissues to wipe the transducer delay line, as they may scratch and damage its surface.



MECHANICAL HAZARD: Ethanol solutions MUST NOT BE USED to disinfect the Bindex device. Ethanol may weaken the plastic cover over time, resulting in loss of mechanical integrity.

In case of visible damage, please contact your local distributor or Bindex Support and Service before attempting to use the Bindex device.

To clean the Bindex BI-2 device and the BI-41 Measure:

1. Wipe impurities, gel residues and other excess matter off the surface of the transducer, the handpiece and the cable as well as the **BI-41** Measure with a dry soft cloth or a soft tissue.

2. A moistened cloth or tissue can be used in case of stains that are harder to remove.

3. Visually inspect the surface of the Bindex BI-2 device and the BI-41 Measure and verify cleanliness.

To disinfect the Bindex BI-2 device and the BI-41 Measure (after cleaning): 1. Use a soft cloth lightly dampened in disinfecting solution to wipe the surface of the transducer and the handpiece as well as the BI-41 Measure. 2. Allow the disinfected parts to dry before using the system again.

After use the Bindex BI-2 device and the BI-41 Measure should be cleaned, disinfected and packed in its protective case to keep them in good condition for the entire lifetime of the system. The packing needs to be done so that no excessive stress accumulates on the device or the USB cord. Make sure that the cord is not sharply twisted or bent and that it is not caught between the edges of the case while closing it.

10. Bindex service

Do not try to service or repair the Bindex device by yourself. There are no selfserviceable or replaceable parts. The Bindex BI-2 device does not require regular servicing by Bone Index Finland. If the device is in need of service or repair (e.g. visible mechanical damage or continuously failing calibration without an apparent reason), please contact your local distributor or Bindex Support and Service (see section 12 for contact details). The expected service life for the Bindex BI-2 device and the BI-41 Measure is three years or 7200 measurements (2400 measurements a year with an average of 10 patient measurements per day, 20 days in month and 12 months).

It is recommended to regularly inspect the condition of you Bindex device, including the casing, the USB cord and the surface of the transducer delay line. The condition of the device should be visually checked every week or at least monthly, depending on the frequency and number of measurements. If the device suffers significant mechanical stress (e.g. falls off a table), its use must be stopped immediately and the condition of the device must be checked. If you notice any changes in the integrity of the device or loss of functionality, please contact your local distributor or Bindex Support and Service before using the device.

The label by the USB connector needs to be legible at all times. The condition of the label should be regularly (recommended monthly) checked to ensure its good condition and legibility. The label is designed to withstand the same disinfection procedures as the device itself so it does not required special care during the cleaning phase. The label MUST NOT BE REMOVED from the device or any accessories. Should the label become illegible or break loose from the device, it needs to be replaced by Bone Index Finland. Please contact your local distributor Bindex Support and Service for further guidance.

If any serious patient/operator incident (death or serious injury) occurs in relation to use of the Bindex device or Bindex Software, the incident must be reported to Bone Index Finland and the competent supervising authority for medical devices in your country without delay.

11. Storing of Bindex

Do not store your Bindex BI-2 in direct sunlight. Sunlight may damage the material properties of the transducer. Store your Bindex in its own case in a dry location and in room temperature.

A quality phantom measurement conducted by Bone Index Finland Ltd. is recommended after one year of continuous storage time before patient measurements.

See section 5 for operating and storing conditions.

11.1. Disposal

Dispose the Bindex BI-2 device according to national or local laws and regulations or according to your disposal policy of your facility regarding Waste Electrical and Electronic Equipment (WEEE).

12. Contact information

Please use your local distributor as a primary contact. If no distributor has been declared, please contact Bindex Support and Service.

General contact address:

Bone Index Finland Oy Savilahdentie 14, 70700 Kuopio, FINLAND Tel. Email: +358 50 448 1696 info@boneindex.fi

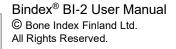
Bindex Support and Service:

Email: info@boneindex.fi Phone: + 358 50 448 1696



INFORMATION SECURITY HAZARD: WARNING: Do not send sensitive information, such as patient health information through email.

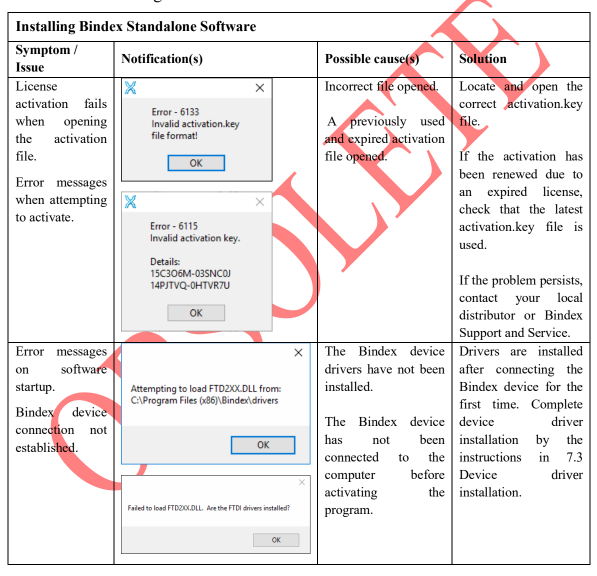
INFORMATION SECURITY HAZARD: WARNING: Verify the email recipient carefully before sending files.



13. Troubleshooting

In case of errors or other issues with Bindex Standalone Software, look for any error messages or notifications. Write down or take a screenshot of any messages and error codes with their details. This will help in resolving possible issues if you need to contact Bindex Support and Service.

Please do not reinstall or delete/modify any files related to the Bindex installation yourself, as this may lead in the activation, patient data and/or activated PPAs being lost.



Using Bindex St	tandalone Software		
Symptom / Issue	Notification(s)	Possible cause(s)	Solution
The measurement cannot be restarted after going to the Results tab.	Proximal tibia measurement is locked!	The measurement is locked after visiting the Results tab. You can go back to the Measurement tab but the measurement cannot be restarted.	To measure the same patient again go to the Main Menu, select OPEN CASE, search and select the patient and click OPEN. Proceed with the measurement as usual.
Unexplained error message(s) when using Bindex Standalone Software.	Examples: Fror 1 occurred at "NI_Database_API.lvlib:Cmd Delete.vi" Possible reason(s): Object 0x000000 is not valid. Continue Stop Stop Fror 0 occurred at an unidentified location Possible reason(s): LabVIEW: Error connecting to GPIB driver or device. UISA: (Hex 0x0) Operation completed successfully. Continue	General error message, may be caused by a number of reasons, e.g. unsuccessful install, insufficient user rights, changes in program files. Always needs to be investigated separately.	In case of a general error situation, write down the: - error number, - location, and - possible reason(s) and send them to Bindex Support and Service. Alternatively, a screen capture of the error message can be taken and sent. Explain where and how the problem occurred. After an error message restart Bindex Standalone Software to prevent incorrect functionality.
Software fails to start. Error message 214767259 on software startup.	 Fror -2147467259 occurred at NLDatabase APLIvlib:DB Tools Open Connec (String).vi >Log - check version and update.vi- > bi- check version, copy and update.vi- >Bindex.vi Possible reason(s): MOD Error: 0x80004005 Exception occurred in Microsoft Access Database Engine: Could not use 'C:\Program Files (x80)\Bindex\log.accdb'; file already in use. in NLD Database_APLI/wlib:DB Tools Open Connec (String).vi->Log - check version and update.vi- >Db - check version, copy and update.vi->Bindex.vi Continue 	Bindex Standalone Software is currently running on another user profile on the computer. The database cannot be used simultaneously by multiple users.	Ask the other user or the system administrator to close Bindex Standalone Software on the other user profile. After this, restart the program.

Symptom / Issue	Notification(s)	Possible cause(s)	Solution
Software fails to start. Error message 7 on startup.	 Error 7 occurred at Move in Db - check version, copy and update.vi->Bindex.vi Possible reason(s): LabVIEW: File not found. The file might have been moved or deleted, or the file path might be incorrectly formatted for the operating system. For example, use \ as path separators on Windows; or MAc OSX, and / on Linux. Verify that the path is correct using the command prompt or file explorer. C:\Program Files (x86)\Bindex\database\database.accdb [6139@] Continue 	The database file is not found, possible reasons: The database location is incorrectly defined in "settings.ini". The database file has been deleted, moved or renamed so that the program cannot find it.	Users with sufficient understanding of file editing: open "settings.ini" in Notepad and check the line "database_directory". Then verify that "database.accdb" file is found in the folder defined on the line. Correct the path, if needed. If unsure or the issue cannot be clearly identified, contact Bindex Support and Service for assistance.
Error message when editing software Settings.	X Settings file is missing. Changes in settings can't be saved and won't affect. Restart software to correct this problem. OK	The Bindex settings file is not found. The "settings.ini" file has been deleted, moved or renamed during program use.	Relocate or rename the "settings.ini" file, if possible. If the original file has been lost, restart software. A new settings file is generated. The user and organizational data must be re-entered in the Settings page. If the problem persists, contact your distributor or Bindex Support and Service.
User Manual cannot be opened in software.	X User manual file not found.	The User Manual file has been moved, renamed or deleted from its original location. The User Manual files are installed in the "user manual" folder of the Bindex install.	At least the default manual in English should be found in the folder with the name "user manual_EN.pdf". Relocate or rename the User Manual file, if possible If the file is missing, please contact Bindex Support and Service for assistance.

Symptom / Issue	Notification(s)	Possible cause(s)	Solution
Calibration of the Bindex device fails.	Calibration failed, please make sure that transducer head is clean and calibrate again. OK Calibration failed! Bindex device might be too cold. Please let the device warm up before use. (p-0,093-t-21,884-c-27,0) OK	The transducer surface is not clean. The storing or operating temperature of the Bindex device is or has been outside of set limits (see the product labels). The transducer is damaged.	the transducer. Software measures the echo signal from the surface of the transducer, which must be clean from any gel or other

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Appendix: Guidance and manufacturer's declaration - Electromagnetic Compatibility

Bindex is intended for use in an electromagnetic environment as specified in section 6 Operating environment. The customer or the user of the Bindex should assure that it is used in such an environment.

Essential performance is defined as maintaining the manufacturer defined accuracy for the apparent thickness measurement even during tested electromagnetic disturbances. Possible degradation or loss of performance due to electromagnetic disturbances can appear as one or more of the following:

- 1) inability to establish or maintain connection to the measurement PC,
- 2) unusual, possibly regular or repeating disturbances or patterns in the signal,
- 3) Bindex software unable to accept the measured signal due to disturbances,
- 4) incorrectly accepted signals with disturbances.

Electromagnetic compatibility along EN 60601-1-2:2015 applies only for an unmodified Bindex BI-2 device with its original cable 9355 installed by Bone Index Finland Ltd. There are no user-replaceable parts or components.

ELECTROMAGNETIC EMISSIONS		
Emission test	Compliance	Environment guidance
RF emissions CISPR 11	Group 1	Bindex uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class B	Bindex should be operated indoors, in clinics, hospitals or home
Harmonic emissions IEC 61000-3-2	Not applicable	healthcare use environments.
Voltage fluctuations/flicker emissions IEC 61000-3-3	Not applicable	

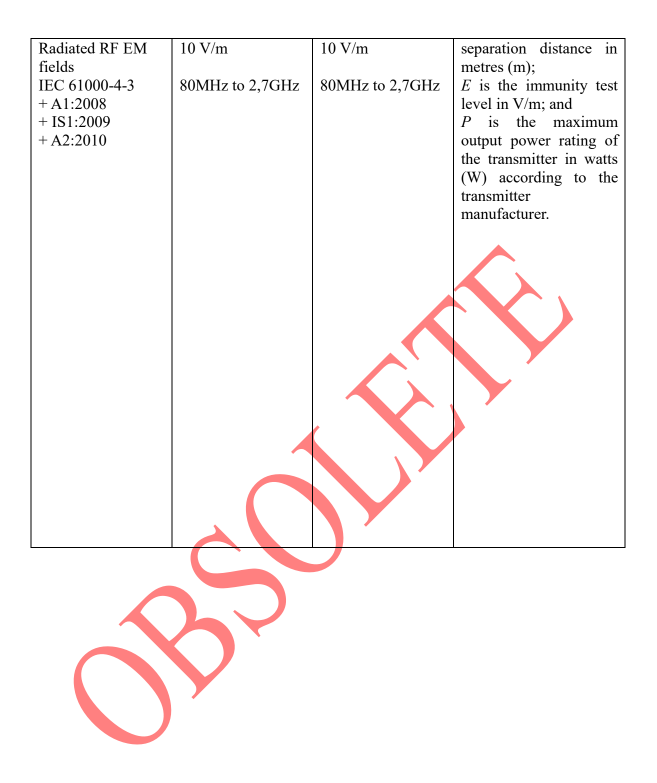
Table 1: Guidance and manufacturer's declaration – Electromagnetic Emissions.

ELECTROMAGNETIC IMMUNITY			
Immunity test	IEC 60601	Compliance	Environment guidance
	test level	level	
Electrostatic	±8kV contact	±8kV contact	Floors should be wood,
discharge (ESD)	±15kV air	±15kV air	concrete or ceramic tile. If
IEC 61000-4-2			floors are covered with
			synthetic material, the relative
			humidity should be at least
			30 %.
Electrical fast	$\pm 2kV$	Not	Mains power quality should be
transient/burst	100 kHz	applicable	that of a typical home
IEC 61000-4-4	repetition		healthcare use, commercial or
	frequency		hospital environment.
Surge	±0,5 kV,	Not	Mains power quality should be
IEC 61000-4-5	±1 kV	applicable	that of a typical home
	Line-to-line		healthcare use, commercial or
	+0.5.1-37		hospital environment.
	±0,5 kV, ±1 kV,		
	$\pm 1 \text{ kV},$ $\pm 2 \text{ kV}$		
	$\pm 2 \text{ KV}$ Line-to-		
	ground		
Voltage dips, short	$0 \% U_T; 0,5$	Not	Mains power quality should be
interruptions and	cycle	applicable	that of a typical home
voltage variations	At 0° , 45° ,	upphonoic	healthcare use, commercial or
on power supply	90°, 135°,		hospital environment. If the
input	180°, 225°,		user of Bindex requires
lines	270° and		continued operation during
IEC 61000-4-11	315°		power mains interruptions, it is
			recommended that Bindex is
	$0 \% U_{T}; 1$		powered from an
	cycle		uninterruptible power supply
	and		or a battery.
	70 % U _T ;		
	25/30 cycles		
	Single phase:		
	at 0°		-
RATED power	30 A/m	30 A/m	Power frequency magnetic
frequency magnetic			fields should be at levels
field	50 Hz or 60		21
IEC 61000-4-8	Hz	Hz	location in a typical home
			healthcare use, commercial or
			hospital environment.
NOTE U _T is the AC mains voltage prior to application of the test level.			

Table 2: Guidance and manufacturer's declaration – Electromagnetic Immunity

ELECTROMAGNETIC IMMUNITY				
Bindex is intended	Bindex is intended for use in the electromagnetic environment specified below. The			
customer or the user	customer or the user of the Bindex should assure that it is used in such an environment.			
Immunity test	IEC 60601 test	Compliance level	Environment	
	level		guidance	
Conducted	3 Vrms	3 Vrms	Field strengths from	
disturbances	0,15 MHz - 80	0,15 MHz - 80	fixed RF transmitters,	
induced by RF	MHz	MHz	as determined by an	
fields			electromagnetic site	
IEC 61000-4-6	6 Vrms in ISM and	6 Vrms in ISM and	survey ^a , should be less	
	amateur bands	amateur bands	than the compliance	
	between 0,15 MHz	between 0,15 MHz	level in each frequency	
	and 80 MHz	and 80 MHz	range. Interference may	
	80 % AM at 1 kHz	80 % AM at 1 kHz	occur in the vicinity of	
			equipment marked with	
		The ISM (industrial,	the following symbol:	
		scientific and medical) bands between 0,15	11	
		MHz and 80 MHz are:	(((•)))	
		6,765 to 6,795 MHz	NA 7	
		13,553 to 13,567 MHz	—	
		26,957 to 27,283 MHz	The minimum	
		40,66 to 40,70 MHz	separation distance for	
		The amateur radio	portable RF	
		bands between 0,15	communications	
		MHz and 80 MHz are:	devices and the Bindex	
		1,8 to 2,0 MHz	device can be calculated	
		3,5 to 4,0 MHz 5,3 to 5,4 MHz	as follows.	
		7 to 7,3 MHz		
		10,1 to 10,15 MHz	Minimum separation	
		14 to 14,2 MHz	distance	
		18,07 to 18,17 MHz		
		21,0 to 21,4 MHz 24,89 to 24,99 MHz	$d = \frac{6}{F}\sqrt{P}$	
		28,0 to 29,7 MHz	u - E v r	
		50,0 to 54,0 MHz		
			where d is the minimum	

Table 3: Guidance and manufacturer's declaration – Electromagnetic Immunity



Proximity fields	9 V/m, 27 V/m, 28	9 V/m, 27 V/m, 28
from RF wireless	V/m	V/m
communications		
equipment	80 MHz to 5800	80 MHz to 5800
EN 61000-4-3	MHz	MHz
+A1:2008		
+ IS1:2009	380 MHz to 390 MHz:	380 MHz to 390 MHz:
+ A2:2010	27 V/m	27 V/m
	430 MHz to 470 MHz:	430 MHz to 470 MHz:
	28 V/m	28 V/m
	704 MHz to 787 MHz: 9 V/m	704 MHz to 787 MHz: 9 V/m
	y thin	
	800 MHz to 960 MHz:	800 MHz to 960 MHz:
	28 V/m	28 V/m
	1700 MHz to 1990 MHz:	1700 MHz to 1990 MHz:
	28 V/m	28 V/m
	2400 MHz to 2570 MHz:	2400 MHz to 2570 MHz:
	28 V/m	28 V/m
	5100 MIL (5900 MIL	
	5100 MHz to 5800 MHz: 9 V/m	5100 MHz to 5800 MHz: 9 V/m
NOTE 1 A + 90 MIL	and 200 MILE the Co	

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies. NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which Bindex is used exceeds the applicable RF compliance level above, Bindex should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating Bindex.